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GenCore version 5.1.6
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Run on:

June 3, 2005, 06:53:47; Search time 17 Seconds (without alignments) 50.938 Million cell updates/sec

US-10-623-176-2 45 1 VLHDDLLEA 9

Title: Perfect score: Sequence:

Gapop 10.0 , Gapext 0.5 **BLOSUM62** Scoring table:

283416 seqs, 96216763 residues Searched:

283416 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

PIR_79:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Description	hypothetical prote	otei	othetic		El protein - human	type I restriction	chemotaxis regulat	probable glyceroph		probable glyceroph		hypothetical prote	-	conserved hypothet	probable dehydroge	polyprenyl synthas	octaprenyl-diphosp	DNA polymerase III		DNA polymerase III	DNA-directed DNA p	probable polypreny		probable N-Acylami	unspecific monooxy	unspecific monooxy		probable peptidase	
ID	S51855	D59433	T00151	S36487	S36469	C69226	A81316	A41652	C89075	T44122	E89762	876860	AG0461	G82601	D83371	B75457	C83075	AI0581	B85564	F90713	A45251	T35403	809813	H72660	869988	869989	\$36493	AH0668	69083
DB	2	1 0	7	7	~	~	7	~	7	~	~	7	~	~	7	~	7	7	7	~	~	~	~	~	-	~	~	7	c
Length	1148	1165	119	604	605	616	130	145	174	210	210	238	249	259	291	320	322	343	343	343	343	378	397	398	519	519	604	654	667
* Query Match	84.4			77.8	77.8	77.8	75.6	73.3	73.3	73.3		73.3		73.3		73.3	73.3			73.3	73.3	73.3				73.3		73.3	73.3
Score	38	37	35	35	35	35	34	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33	33
Result No.	-	10	Ю	4	S	9	7	ω	6	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	59

A,Reference number: D59433; MUD:97191544; PMID:9039502
A;Accession: D59433
A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-1165 < NAG>
A;Cross-references: UNIPROT:Q92619; GB:BAA13212; PID:g1504026; PIDN:BAA13212.1
S;Chara, O.; Nagase, T.; Kikuno, R.; Nomura, N.
submitted to Genbank, August 1996
A;Reference number: E59433
A;Accession: E59433
A;Accession: E59433
A;Actus: preliminary
A;Catus: preliminary
A;Catus: Dreliminary
A;Residues: 1-1165 <STO2>

Ä~.	hypothetical proce seck protein (impo DNA repair and rec	MDM1 protein - yea hypothetical prote ribonuclease inhib acidic ribosomal p	A-ORF-E protein hypothetical prote hypothetical prote	probable thiosulfa hypothetical prote probable trans-pre
H85734 F64895	C85023 T50695 T50449	S47445 S56221 C69973 R6KM1C	A42524 G84226 T28746	G70732 T27768 A84759
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667	856 908 973	1127	166 270 276	284 296 297
73.3	7. E. E. 5. E. E. 5. E. E. E.	73.3	71.17	71.1
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330	1 W W W 1 W 4 R	33 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9	4 4 4 2 1 2 2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3	4 4 4 5 4 5

ALIGNMENTS

RESULT S51855	F 1
hypoth N,Alte	hypothetical protein YDR128w - yeast (Saccharomyces cerevisiae) N;Alternate names: hypothetical protein YD9302.03
C;Spec	C;Species: Saccharomyces cerevisiae C:Date: 05-May-1995 #sequence revision 21-Jul-1995 #text change 16-Aug-2004
C, Acce	C.Accession: S51855 E.S. E.S. E.S. E.S. E.S. E.S. E.S. E.
submit A:Refe	submitted to the EMBL Data Library, February 1995 A.Reference number: S51853
A, Acce	A,Accession: SS1855
A; MOLE A; Resi	A;Wolecule type: DNA A;Residues: 1-1148 <oli></oli>
A;Cross-ref	A;Cross-references: UNIPROT:Q03897; EMBL:Z48179; NID:g665657; PID:g665660; GSPDB:GN00004;
A; Gene	Y,GENET MIPS:YDR128w
A;Cros	A,Cross-references: SGD:S0002535
A;Map C;Supe	A,Map position: 4R C,Superfamily: WD repeat homology
Quer	84.48;
Best	Best Local Similarity 66.7%; Pred. No. 52; Matches 6; Conservative 3; Mismatches 0; Indels 0; Gaps 0;
ò	1 VLHDDLLEA 9
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RESULT	2 1
D59433 C. elec	D59433 C. elegans protein Z37093 homolog [imported] - human
C,Spec	C.Species: Homo sapiens (man) C.Date: 01-Jnn-2002 #semience revision 01-Jnn-2002 #text change 09-Jn1-2004
C, Acce	C;Accession: D59433; E59433
R; Naga	R:Nagase, T.; Seki, N.; Ishikawa, K.; Ohira, M.; Kawarabayasi, Y.; Ohara, O.; Tanaka, A.; nna bes 1 121-122
A,Titl	A, Title: Prediction of the coding sequences of unidentified human genes. VI. The coding f

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A;Molecule type: DNA
A;Residues: 1-605 <DEL>
A;Cross-references: UNIPROT:P36721; EMBL:X74467; NID:g396918; PIDN:CAA52502.1; PID:g39691
A;Experimental source: strain 14D
C;Superfamily: papillomavirus El protein
C;Keywords: early protein; nucleus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Cjaccession: C69226
R;Smith, D.R.; Doucette-Stamm, L.A.; Deloughery, C.; Lee, H.; Dubois, J.; Aldredge, T.; F.; Qiu, D.; Spadafora, R.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Gibson, R.; Jiwani, N.; Qiu, D.; Spadafora, R.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Reeve, J.N.
J. Bacteriol. 179, 7135-7155, 1997
A; Title: Complete genome sequence of Methanobacterium thermoautotrophicum Delta H: functi A; Reference number: A69000; MUID:98037514; PMID:9371463
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     chemotaxis regulatory protein Cj1118c [imported] - Campylobacter jejuni (strain NCTC 1116 C;Species: Campylobacter jejuni C;Date: 31-Mar-2000 #sequence_revision 31-Mar-2000 #text_change 16-Aug-2004 C;Accession: A81316
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        R.Parkhill, J.; Wren, B.W.; Mungall, K.; Ketley, J.M.; Churcher, C.; Basham, D.; Chilling C.W.; Quail, M.; Rajandream, M.A.; Rutherford, K.M.; VanVliet, A.; Whitehead, S.; Barrell Nature 403, 665-668, 2000
A;Title: The genome sequence of the food-borne pathogen Campylobacter jejuni reveals hype A;Reference number: A81250; MUID:20150912; PMID:10688204
A;Accession: A81316
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A;Cross-references: UNIPROT:027025; GB:AE000868; GB:AE000666; NID:g2622025; PIDN:AAB8544(
A;Experimental source: strain Delta H
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A,Experimental source: serotype O2, strain NCTC 11168
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           type I restriction modification enzyme, subunit M - Methanobacterium thermoautotrophicum C;Species: Methanobacterium thermoautotrophicum C;Date: 05-Dec-1997 #sequence_revision 05-Dec-1997 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               A;Status: preliminary; nucleic acid sequence not shown; translation not shown A;Wolecule type: DNA
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C;Superfamily: Signal transduction receiver (phosphoacceptor) protein, CheY
                                                                                                                                                                                                                                                                  Gaps
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Pred. No. 96;
1; Mismatches
                                                                                                                                                                                                                  ;;
                                                                                                                                                                                                               Score 35; DB 2; Pred. No. 94; 2; Mismatches
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Local Similarity 75.0%;
nes 6; Conservative
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Best Local Similarity 77.00,
7, Conservative
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Matches 6; Conservative
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202 VHDDLLES 209
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A; Residues: 1-130 <PAR>
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A; Accession: S36469
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A;Residues: 1-604 <DBL:
A;Cross-tences: UNIPROT:Q02048; EMBL:X74470; NID:g396940; PIDN:CAA52520.1; PID:g3969
C;Superfamily: papillomavirus El protein
C;Keywords: early protein; nucleus
                                                                                                                                                                                                                                                                                                                    hypothetical protein 18 - Staphylococcus aureus phage phi PVL
C;Species Staphylococcus aureus phage phi PVL
C;Descies Staphylococcus aureus phage phi PVL
C;Accession: T00151
R;Kaneko, J.; Kimura, T.; Kawakami, Y.; Tomita, T.; Kamio, Y.
Bjoeci. Biotechnol. Biotenem C., 1960-1962, 1997
A;Title: Panton-Valentine leukocidin genes in a phage-like particle isolated from mitomy A;Reference number: Z14119; MUID:98067870; PMID:9404084
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              A;Cross-references: UNIPROT:080057; EMBL:AB009866; NID:g3341907; PIDN:BAA31891.1; PID:g3
C;Superfamily: Staphylococcus aureus phage phi PVL hypothetical protein 18
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C;Date: 20-Feb-1995 #sequence_revision 20-Feb-1995 #text_change 09-Jul-2004
C;Accession: S36487
R;Delius, H; Hoffmann, B.
aubmitted to the EMBL Data Library, August 1993
A;Description: Primer-directed sequencing of human papillomavirus types.
A;Reference number: S36469
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Bl procein - human papillomavirus type 14D
C:Species: human papillomavirus type 14D
C:Date: 09-Dec-1993 #sequence_revision 26-Jul-1996 #text_change 09-Jul-2004
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Pred. No. 94;
2; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match 77.8%; Score 35; DB 2; Length 119; Best Local Similarity 85.7%; Pred. No. 15; Matches 6; Conservative 1; Mismatches 0; Indels
A; Cross-references: GB: BAA13212; PID: g1504026; PIDN: BAA13212.1
                                                        .;
?
                                                  Score 37; DB 2
Pred. No. 82;
0; Mismatches
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                                                     82.2%;
88.9%;
                            Query Match
Best Local Similarity 88.9.
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Best Local Similarity 75.0
Matches 6; Conservative
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166 VLRDDLLEA 174
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201 VHDDLLES 208
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A; Residues: 1-119 < KAN>
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OM protein - protein search, using sw model

Run on:

June 3, 2005, 06:53:02; Search time 81.25 Seconds (without alignments) 56.723 Million cell updates/sec

US-10-623-176-2 45 1 VLHDDLLEA 9

Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

1612378 seqs, 512079187 residues Searched:

1612378 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution. UniProt_03:*
1: uniprot_sprot:*
2: uniprot_trembl:*

	otion	homo sapien	picrophilus	methanococc	saccharomyc	homo sapien	homo sapien	homo sapien	desulfotale	oryza sativ	shewanella	xanthomonas		pseudomonas	mouse adeno	homo sapien	ixodes scap	staphylococ	staphylococ	human papil	human papil	human papil	methanobact	gallus gall	xenopus tro	leptospira	campylobact	geobacter s	anopheles g	streptomyce	gibberella	bacillus sp
	Description	26p189	26kzh5	Q61z81	203897	Q8iyn3	28hx84	092619	26amv9	26aun7	Q8ejs9	Q8pqr0	80057	288cw6	Q7m5u3	26zw54	28mvb1	Q8sdk3	Ogdme0	P50759	202048	P36721	027025	08jhz8	26dir8	Q8eyr5	P71129	Q74b30	07pt12	Q8rjw9	296va9	Q6dv84
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RIES																					•	₹*					JB					
SUMMARIES		189	ZH5	281	168	KN3	X84	619	6Λ <u>Ρ</u>	CN7	389	OR0	057	CW6	5U3	W54	VB1	DK3	BP0	JE1 HPV21	1_HPV19	HPV14	025	HZ8	IR8	YRS	CHEY CAMJE	B30	IL2	6MI	VA9	V84
	£	Q6P189	Q6KZH5	06LZ81	003897	Q81YN3	Q8HX84	092619	Q6AMV9	Q6AUN7	QBEJS9	QBPQR0	080057	Q8SCW6	Q7M5U3	Q6ZW54	QBMVB1	QBSDK3	O9MBP0	Œ	VEI	VE1_HP	027	Q8JHZ8	QEDIRB	QBEYRS	HE	Q74B30	Q7PTL2	Q8RJW9	Q96VA9	Q6DV84
	88	7	7	~	7	~	~	7	N	7	~	N	~	~	~	~	~	~	7	Н	-	-	~	~	~	~		7	7	8	~	7
	Length	1136	571	668	1148	1131	1136	1165	101	231	585	691	119	120	132	148	376	496	496	603	604	605	616	843	872	68	130	146	195	228	241	248
عد	Match	100.0	88.9	86.7	84.4	82.2	82.2	82.2	80.0	80.0	80.0	80.0	77.8	77.8	77.8	77.8	77.8	77.8	77.8	77.8	77.8	77.8	77.8	٠	77.8	75.6	75.6	75.6	75.6	75.6	75.6	75.6
	Score	45	40	39	38	37	37	37	36	36	36	36	35	35	35	35	35	35	35	35	35	35	35	35	35	34	34	. 34	34	34	34	34
	Result No.	1	7	m	4	2	9	7	80	6	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	56	27	28	29	30	31

Q889t7 pseudomonas		O9bjl6 giardia lam			Q87zg8 pseudomonas	Q6k4v7 oryza sativ	Q941f5 oryza sativ	Q84mw4 oryza sativ	Q8nj19 mucor circi	Q8ms72 drosophila	Q7prg2 anopheles g	Q7qsx6 giardia lam	Q9nkm2 leishmania
Q889T7	Q64GC9	Q9BJL6	Q7QYU3	Q65CP2	Q87ZG8	Q6K4V7	Q94LF5	Q84MW4	Q8NJ19	Q8MS72	Q7PRG2	Q7QSX6	Q9NKM2
2	~	3	~	2	7	~	2	2	~	7	7	7	7
253	26.	56€	29.	366	372	401	408	408	47	485	525	536	64(
9.	9.	9.	9.	9.	9.	9.	9.	9.	9.	9.	9.	9.	9.
75.6	75	75	75	75	75	75	75	75	75	75	75	75	75
34	34	34	34	34	34	34	34	34	34	34	34	34	34
32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

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[1]
SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          NCBI_TaxID=4932;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  STRAIN=AB972;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             STRAIN=AB972
                                                                                                                                                                                                                                                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                  003897
                                                                                                                                                                                                                                                                                                                                 Matches
                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 4
Q03897
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                                                                                                                                                                                                                                                                                                                                                                 Fuetterer O., Angelov A., Liesegang H., Gottschalk G., Schleper C., Schepers B., Dock C., Antranikian G., Liebl W.; "Genome sequence of Picrophilus torridus and its implications for life
                                                                                                   Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ö
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                                                                                                                                                                                                                                                                                                                                                                                                         Proc. Natl. Acad. Sci. U.S.A. 101:9091-9096 (2004).

EMBL, AE017261; AAT43877.1; -
GO; GO:0006777; P:Mo-molybdopterin cofactor biosynthesis; IEA.

InterPro; IPR001453; MocA.

InterPro; IPR005110; MocA.

InterPro; IPR005110; MocA.
                                                                            100.0%; Score 45; DB 2; Length 1136;
                                                                                                                                                                                                                                                                                    Archaea; Euryarchaeota; Thermoplasmata; Thermoplasmatales;
Picrophilaceae; Picrophilus.
NCBI_TaxID=82076;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                88.9%; Score 40; DB 2; Length 571; 88.9%; Pred. No. 45; tive 0; Mismatches 1; Indels
                                                                                                   0; Indels
                                                      1136 AA; 124550 MW; 14B7073296102DD5 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Complete proteome.
SEQUENCE 571 AA; 64707 MW; DZEBIBFBFC115FA1 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Methanococcus maripaludis.
Archaes, Buryarchaeots, Methanococci; Methanococcales,
Methanococcaceae; Methanococcus.
NCBI_TaxID=39152;
                                                                                                                                                                                                                      Created)
Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Last sequence update)
Last annotation update)
SMART; SM00324; RhoGAP; 1.

PROSITE; PS00190; CYTOCHROME C; UNKNOWN 1.

PROSITE; PS00479; DAG PE BIND DOM 1; UNKNOWN 1.

PROSITE; PS500081; DAG PE BIND DOM 2; 1.

PROSITE; PS50238; RHOGAP; 1.

SEQUENCE 1136 AA; 124550 WW; 14B7073296102D
                                                                                                                                                                                                                                                                                                                                           STRAIN-DSM 9790 / ATCC 700027;
PubMed=15184674; DOI=10.1073/pnas.0401356101;
                                                                                      9.9;
                                                                                                                                                                                                    571 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        668 AA.
                                                                                                   0; Mismatches
                                                                                                                                                                                                                      05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last sequency
05-JUL-2004 (TrEMBLrel. 27, Last annotat.
Molybdopterin biosynthesis MoeA protein.
OrderedLocusNames=PTO1292;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           05-JUL-2004 (TrEMBLrel. 27, Created)
05-JUL-2004 (TrEMBLrel. 27, Last seq
05-JUL-2004 (TrEMBLrel. 27, Last ann
MCM family related protein.
OrderedLocusNames=MMP0748;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          ProDom; PD002460; MoCF biosynth; 1.
TIGRFAMB; TIGR00177; molyb_syn; 1.
                                                                                                                                                                                                   PRT;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Pfam; PF00994; MoCF biosynth; 1. Pfam; PF03454; MocA_C; 1. Pfam; PF03453; MocA_N; 1.
                                                                                        100.0%;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ilarity 88.9%;
Conservative
                                                                                       Best Local Similarity 100.
Matches 9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       PRELIMINARY;
                                                                                                                                                                                                   PRELIMINARY;
                                                                                                                                       137 VLHDDLLEA 145
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 VLHDDLLNA 71
                                                                                                                                                                                                                                                                              Picrophilus torridus.
                                                                                                                        1 VLHDDLLEA 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             1 VLHDDLLEA 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Local Similarity
Les 8; Conserv
                                                                                                                                                                                                                                                                                                                                  SEQUENCE FROM N.A
                                                                                                                                                                                                                                                                                                                                                                                                  around pH 0."
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
                                                                             Query Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Q6LZ81
Q6LZ81;
                                                                                                                                                                                                 Q6KZH5
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Loc
Matches
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QGKZH5
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STRAIN=S2 / LL;
Hendrickson E.L., Kaul R., Zhou Y., Bovee D., Chapman P., Chung J., Conway de Macario E., Dodsworth J., Gillett W., Graham D.E., Haydock A.K., Kang A., Land M.L., Levy R., Lie T.J., Major T., Moore B., Porat I., Overbeek R., Palmeiri A., Rouse G., Saenphimmachak C., Soll D., Whitman W.B., Larimer F.W., Olson M.V.,
                                                                                                                                                                                                       "Complete grown sequence of the mesophilic hydrogenotrophic methanogen Methanococcus maripaludis.";
In Submitted (FEB-2004) to the EMBL/GenBank/DDBJ databases.
In Submitted (FEB-2004) to the EMBL/GenBank/DDBJ databases.

R GO; GO:0005524; F:ATP binding; IEA.

R GO; GO:0008094; F:DNA binding; IEA.

R GO; GO:0008094; F:DNA replication initiation; IEA.

R InterPro; IPR002048; EF-hand.

R InterPro; IPR008094; Nucleic_acid_OB.

R PRINTS; PR01657; MCMPAMILY.

R PRODOM; GO:00041; MCM; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Saccharomyces cerevisiae (Baker's yeast).
Eukaryota; Fungi; Ascomycota; Saccharomycotina; Saccharomycetes;
Saccharomycetales; Saccharomycetaceae; Saccharomyces.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ;
0
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Submitted (FEB-1995) to the EMBL/GenBank/DDBJ databases.
-: -: SIMILARITY: Contains 4 WD repeats.
EMBL; Z48179; CAR88209.1; -.
PIR; S51855; S51855.
SGD; S00000253; YDR128W.
GO; GO:0000253; YDR128W.
GO; GO:0000329; C:vacuolar membrane (sensu Fungi); IDA.
InterPro; IPR006575; RWD.
InterPro; IPR00400; WD40; -.
PRINTS; PR00320; GPROTEINBRPT.
SMART; SM00320; GPROTEINBRPT.
SMART; SM00320; WD40; -.
PROSITE; PS500678; WD.
PROSITE; PS500678; WD.
PROSITE; PS00678; WD.
PROSITE; PS00678; WD.
PROSITE; PS00678; WD. REPEATS_1; UNKNOWN_1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Length 668
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Oliver K., Harris D.;
Submitted (FEB-1995) to the EMBL/GenBank/DDBJ databases
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  668 AA; 75660 MW; 2ADAF3800B1049F4 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Q03897;
01-NOV-1996 (TrEMBLrel. 01, Created)
01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-NAX-2004 (TrEMBLrel. 26, Last annotation update)
Hypothetical protein.
ORFNames=YDR128W;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          86.7%; Score 39; DB 2;
88.9%; Pred. No. 82;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PRT; 1148 AA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           1; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   PROSITE; PS00018; EF HAND; UNKNOWN_1.
PROSITE; PS50051; MCM_2; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              8; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             ||:||||||
628 VLYDDLLEA 636
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SMART; SM00350; MCM; 1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1 VLHDDLLEA 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Complete proteome.
SEQUENCE 668 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Best Local Similarity
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	Ltd.
5.1.6	Compugen
version	- 2005
GenCore	(c) 1993
	Copyright

3, 2005, 06:44:47; Search time 90.5 Seconds June Run on:

(without alignments) 38.462 Million cell updates/sec

US-10-623-176-2 45 Title:

1 VLHDDLLEA 9 Perfect score: Sequence:

BLOSUM62 Scoring table:

Gapop 10.0 , Gapext 0.5

2105692 seqs, 386760381 residues Searched:

2105692 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100

Listing first 45 summaries Match 100%

A_Geneseq_16Dec04:* 1: qeneseqp1980s:*

Database

geneseqp1980s:* geneseqp1990s:*

geneseqp2003as:* geneseqp2003bs:* geneseqp2000s:* geneseqp2001s:* geneseqp2002s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

geneseqp2004s:*

SUMMARIES

		de			SUMMAKIES		
	Score	Query Match	Length	DB	aı	Description	
-	45	100.0	6	7	AAW99196	Aaw99196 Minor his	
7	45	100.0	Q	N	AAW97375	Aaw97375 HA-1 H-al	
٣	45	100.0	Ø.	æ	ADH40333	Adh40333 Human min	
4	45	100.0	13	~	AAW99199	σ	
ß	45	100.0	13	~	AAW97415	Aaw97415 KIAA0223	
9	45	100.0	1165	æ	ADS11768	Ads11768 Human the	
7	38	84.4	199	7	ADF06830	Adf06830 Bacterial	
œ	37	82.2	o,	~	AAW99197	Aaw99197 Minor his	
σ	37	82.2	0	~	AAW97374	Aaw97374 HA-1 R-al	
10	37	82.2	Φ	œ	ADH40334	Adh40334 Human min	
11	37	82.2	13	~	AAW99198	Aaw99198 DH cell K	
12	37	82.2	13	~	AAW97414	Aaw97414 KIAA0223	
13	37	82.2	1136	'n	AD117193	Adil7193 Human NOV	
14	37	82.2	1136	Ŋ	ADI16744	Adil6744 Human NOV	
15	37	82.2	1136	œ	ADH61299	Adh61299 INTSIG pr	
16	37	82.2	1136	œ	ADN42398	Adn42398 Human nov	
17	37	82.2	1136	œ	ADS10490	Ads10490 Human the	
18	37	82.2	1165	'n	ADI17194	Adil7194 Human NOV	
19	37	82.2	1165	œ	ADP55035	Adp55035 Human PRO	
20	37	82.2	1165	80	ADP24600	Adp24600 PRO polyp	
21	36	80.0	6	7	AAW99195		
22	36	80.0	o,	~	AAW97572	Aaw97572 T-cell ep	
23	35	77.8	148	œ	ADQ65728	Add65728 Novel hum	
24	35	77.8	496	9	ABM71411	Abm71411 Staphyloc	
25	35	77.8	609	7	AB077464	Abo77464 Pseudomon	

Abu29875 Protein e Adc97107 E. faeciu	Pro				Adj48425 Maize oil Abb65640 Drosophil	Aau38963 Drosophil Adc35871 Drosophil	Aag91263 C glutami Aae38199 Fruit fly	Abb63586 Drosophil	
ABU29875 ADC97107	ABU26589	ABM52112 ABU42064	ABB64157 ABU41832	ABD14623 ABU34950	3 ADJ48425 3 ABB65640	AAU38963 ADC35871	AAG91263	ABB63586	ABU31332
786 6	130 6	232 6	332 385 6	440 446 6	680 774 4	774 477	783 4	1383 4	124 6
77.8	75.6	75.6	75.6 75.6	75.6	75.6	75.6 75.6	75.6	75.6	73.3
33.5	0 W W	6. E	344	344	34 34	34	34	34	33
27	300	31	а а 4	32 36	37 38	39 40	41	64.	45

ALIGNMENTS

RESULT 1 AAW99196

AAW99196 standard; peptide; 9 AA.

AAW99196;

(first entry) 20-MAY-1999 Minor histocompatibility antigen HA-1 T-cell epitope #2.

Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.

Homo sapiens

WO9905174-A1

04-FEB-1999.

98WO-NL000425. 23-JUL-1998; 97EP-00202303. 23-JUL-1997;

(UYLE-) RIJKSUNIV LEIDEN.

Engelhard VH; Hunt DF, Goulmy EAJM,

WPI; 1999-153312/13.

A new minor histocompatibility antigen, HA-1 - useful to treat immune diseases and prevent rejection and host versus graft disease in bone marrow and organ transplantation.

Claim 3; Page 32; 47pp; English.

epitope obtainable from the minor histocompatibility antigen HA-1. The peptide is immunogenic and can be used as part of a vaccine. Pl is used as a medicine, to induce tolerance for transplants, prevent rejection and/or graft versus host disease, or to treat (auto) immune diseases. In particular it can be used with bone marrow transplantation, in the treatment of severe aplastic anaemia, leukaemia, and immune deficiency The present sequence represents a new peptide (P1) constituting a T-cell

Sequence 9 AA;

Query Match

100.0%; Score 45; DB 2; Length 9;

Matches

RESULT 2

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The invention relates to a novel method for providing epitopes of allelic variants of antigenic proteins from specific species based on single nucleotide polymorphism (SNP), by defining target protein/peptide or its subset, screening database of DNA encoding target protein, identifying, selecting allelic peptide/protein variants, expression product or its fragment encoded by DNA sequence having SNP, creating variant epitopes, conditing epitopes binding to MC protein. A protein of the invention has cytostatic activity, and may have a use in a vaccine. The method is useful for generating a SNP profile of one or more individuals from a given species by applying the method for several protein from the individuals, where the SNP profile was related to disease, preferably cancer. This is useful for diagnoshing a disease in an individual by cancer. This is useful for diagnoshing a disease in an individual by cancer transplanting haematopoietic stem cells from a donor to a recipient and treating cancer, preferably leukaemia, and for determining the progression, regression or onset of a treated disease. The present sequence is used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Providing allelic variant epitope of protein based on single nucleotide polymorphism by defining target protein, screening database of protein, identifying, selecting allelic variant protein, creating variant
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.
                                                                                    human, cytostatic, vaccine, SNP profile; cancer, leukaemia, minor histocomptibility antigen, mHAg; T cell epitope.
                                          Human minor histocomptibility antigen HA-1 T cell eptiope.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             100.0%; Score 45; DB 8; I
Similarity 100.0%; Pred. No. 1.8e+06;
9; Conservative 0; Mismatches 0;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Disclosure, Page 82; 119pp; English
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     vR cell KIAA0223 protein sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AAW99199 standard; peptide; 13 AA.
                                                                                                                                                                                                                                                                                                                                         13-JUN-2002; 2002EP-00013423.
                                                                                                                                                                                                                                                                                               13-JUN-2003; 2003WO-EP006251.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        (first entry)
11-MAR-2004 (first entry)
                                                                                                                                                                                                                                                                                                                                                                                   (MERE ) MERCK PATENT GMBH.
                                                                                                                                                                                                                                                                                                                                                                                                                               Strittmatter W, Moll H;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          σ
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                                                                                                                                                                                                      WO2003106692-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Sequence 9 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        20-MAY-1999
                                                                                                                                                              Homo sapiens
                                                                                                                                                                                                                                                  24-DEC-2003.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAW99199;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
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                                                                                                                                                                                                                                                                                                                                                                                                        Intron; minor histocompatibility antigen HA-1; typing allele; H allele; R allele; polymorphic nucleotide; HA-1 typing; bone marrow transplant; severe aplastic anaemia; leukaemia; immune deficiency disease; ss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Typing minor histocompatibility antigen HA-1 - by amplifying and identifying HA-1 R and HA-1 H alleles of HA-1, used for, e.g. detection
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Local Similarity 100.0%; Pred. No. 1.8e+06;
Heb 9; Conservative 0; Mismatches 0;
  Pred. No. 1.8e+06;
                         0; Mismatches
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Gaps

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RESULT 3 ADH40333

Length 9; Indels

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Fri Jun 3 0
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
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1: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/Ptr_NEW_PUB_Dep:*} \)

2: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/Ptr_NEW_PUB_Dep:*} \)

3: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/NSOG_NEW_PUB_Dep:*} \)

4: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOG_NEW_PUB_Dep:*} \)

5: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOG_NEW_PUB_Dep:*} \)

6: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOG_NEW_PUB_Dep:*} \)

7: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOB_NEW_PUB_Dep:*} \)

9: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOB_NEW_PUB_Dep:*} \)

10: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOB_PUBCOMB_Dep:*} \)

11: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOB_PUBCOMB_Dep:*} \)

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14: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USOB_PUBCOMB_Dep:*} \)

15: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_PUBCOMB_Dep:*} \)

16: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_PUBCOMB_Dep:*} \)

17: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_PUBCOMB_Dep:*} \)

18: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_PUBCOMB_Dep:*} \)

19: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_PUBCOMB_Dep:*} \)

17: \cgn2 \( \bigcore{G}\) \text{ptodata/1/pubpaa/USID_NEW_PUB_Dep:*} \)

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17: \cgn2 \( \bigcore{G}\) \t
GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-10-791-217-2
US-10-623-176-43
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ALIGNMENTS

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Sequence 2, Application US/10623176

Sequence 2, Application No US20040092446A1

GENERAL INCOPMATION:

APPLICANT: Goulmy, Els A.J.M.

APPLICANT: Goulmy, Els A.J.M.

APPLICANT: Hunt, Donald F.

APPLICANT: Engelhard, Victor H.

TITLE OF INVENTION: HA.1 epitopes and uses thereof

FILE REFERENCE: 2183-60470S

CURRENT FILING DATE: 2000-01-21

PRIOR APPLICATION NUMBER: 09/489,760

PRIOR APPLICATION NUMBER: P97202303.0

PRIOR APPLICATION NUMBER: P97202303.0

PRIOR PILING DATE: 1998-07-23

PRIOR PILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

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OTHER INFORMATION: sequence derived from a presumed HA-1 homozygous
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                                                        APPLICANT: Goulmy, Bls A.J.M.
APPLICANT: Goulmy, Bls A.J.M.
APPLICANT: Goulmy, Bls A.J.M.
APPLICANT: Hunt, Donald F.
TITLE OF INVEXTION: HA-1 epitopes and uses thereof
FILE REFERENCE: 2183-60470S
CURRENT APPLICATION NUMBER: US/10/623,176
CURRENT PLILNG DATE: 2003-07-18
PRIOR APPLICATION NUMBER: 09/489,760
PRIOR APPLICATION NUMBER: P97202303.0
PRIOR APPLICATION NUMBER: P770-23
PRIOR APPLICATION NUMBER: P770-23
PRIOR PILING DATE: 1998-07-23
PRIOR FILING DATE: 1998-07-23
PRIOR FILING DATE: 1998-07-23
PRIOR FILING DATE: 2000-01-24
NUMBER OF SEQ ID NOS: 77
SOFTWARE: Patentin Ver. 2.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence 76, Application US/10623176 Publication No. US20040092446A1 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              TYPE: PRT
ORGANISM: Artificial Sequence
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ORGANISM: Artificial Sequence
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Best Local Similarity
Matches 9; Conserv
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US-10-623-176-76
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US-10-623-176-43
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; Sequence 1, Application US/10861335
; Publication No. US20050031612A1
; GENERAL INFORMATION:
; APPLICANT: Goulmy, Elsa A.J.M.
; TITLE OF INVENTION: Minor histocompatibility antigen HA-1: target antigen for immunot; TITLE OF INVENTION: Minor histocompatibility antigen HA-1: target antigen for immunot; TITLE OF INVENTION: Minor histocompatibility antigen HA-1: target antigen for immunot; TITLE OF INVENTION: Minor histocompatibility antigen HA-1: target antigen for immunot; TITLE OF INVENTION: 04 tunner: 2004-06-04
; FILE REFRENCE: 2004-06-04
; PRIOR APPLICATION NUMBER: EP 01204-06-04
; PRIOR FILING DATE: 2001-12-05
; SOFTWARE: PAPLICATION NUMBER: EP 01204704.9
; SOFTWARE: PAPLICATION NUMBER: BP 01204704.9
; SOFTWARE: PALENCE PAPLICATION OF 10.05
; SOFTWARE: PAPLICATION OF 10.05
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           0; Mismatches
                                                                                                                                                                                                                            APPLICANT: Goulmy, Elsa A.J.M
APPLICANT: Goulmy, Elsa A.J.M
APPLICANT: Hunt, Donald F
APPLICANT: Bigelhard, Victor H
TITLE OF INVENTION: The HA-1 Antigen
FILE REFERENCE: 2183-4285US
CURRENT APPLICATION NUMBER: US/10/791,217
CURRENT APPLICATION NUMBER: US/09/489,760
PRIOR FILING DATE: 2000-01-21
PRIOR FILING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: Patentin Ver. 2.1
SEG ID NO
                                                                                                                                                              Sequence 2, Application US/10791217
Publication No. US20040191268A1
GENERAL INFORMATION:
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US-10-861-335-1
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ORGANISM: Artificial Sequence
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           9; Conservative
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1 VLHDDLLEA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   TYPE: PRT
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Sequence 7115, Ap
Sequence 156, App
Sequence 26, Appl
Sequence 229, Appl
Sequence 2314, Ap
Sequence 31480, A
Sequence 3701, Ap
Sequence 2701, Ap
Sequence 2701, Ap
Sequence 27129, A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                 3, 2005, 06:54:48 ; Search time 24 Seconds (without alignments) 27.993 Million cell updates/sec
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/cgm2_6/ptodata/1/iaa/5B_COMB.pep:*
/cgm2_6/ptodata/1/iaa/6A_COMB.pep:*
/cgm2_6/ptodata/1/iaa/6B_COMB.pep:*
/cgm2_6/ptodata/1/iaa/PcTUS_COMB.pep:*
/cgm2_6/ptodata/1/iaa/PcTUS_COMB.pep:*
GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-269-250R-28
US-09-269-250R-38
US-09-543-681A-7115
US-09-269-250R-18
US-09-269-250R-18
US-09-269-250R-29
US-09-269-250R-29
US-09-275-991A-26210
US-09-107-532A-3480
US-09-270-767-48697
US-09-134-000C-3701
US-09-134-000C-4544
US-09-134-001C-4544
US-09-134-001C-4544
US-09-252-991A-2148
US-09-818-708-97
US-09-818-708-97
US-09-818-708-97
US-09-818-708-97
US-09-818-708-97
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US-09-489-039A-9868
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US-09-071-035-334
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                                                                                                                                                                                                                                                                                                                                                  Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                          OM protein - protein search, using sw model
                                                                                                                                                                                                                    Gapop 10.0 , Gapext 0.5
                                                                                                                                                                                                                                                                                                                                                                                                            Issued_Patents_AA:*
                                                                                                                                                                                                                                                                                                       seq length: 0
seq length: 200000000
                                                                                                                                             US-10-623-176-2
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Match Length DB
                                                                                                                                                                       1 VLHDDLLEA 9
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                                                                                                                                                            Perfect score:
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No.
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Sequence 20, Application US/09269250E
Patent No. 6830883
GENERAL INFORMATION:
GENERAL INFORMATION:
GENERAL INFORMATION:
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REFERENCE: 58994
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ 1D NOS: 38
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Patent No. 6830883
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: Goulmy, Elsa
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REPRENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269, 250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
SEQ ID NO 28
LENGTH: 13
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ; OTHER INFORMATION: Description of Artificial Sequence: Exon Fragments US-09-269-250E-20
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      , OTHER INFORMATION: Description of Artificial Sequence: PCR Product US-09-269-250E-28
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Sequence Seq
       Sequence
Seq
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illarity 100.0%; Pred. No. 4.1e+05;
Conservative 0; Mismatches
US-09-252-991A-26814
US-09-428-177-2
US-09-082-2798-454
US-09-082-2798-455
US-08-474-349A-481
US-08-474-349A-481
US-08-474-349A-481
US-09-315-3048-454
US-09-315-3048-455
US-09-315-3048-455
US-09-314-784-456
US-09-834-784-456
US-09-834-784-456
US-09-834-784-456
US-09-834-784-456
US-09-815-965A-456
US-09-515-965A-456
US-09-515-965A-456
US-09-515-965A-456
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ORGANISM: Artificial Sequence
FEATURE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SOFTWARE: PatentIn version 3.1 SEQ ID NO 20
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           TYPE: PRT
ORGANISM: Artificial Sequence
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Best Local Similarity
9; Conserv?
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       US-09-269-250E-20
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       LENGTH:
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Length 13;

DB 4;

100.0%; Score 45;

Query Match

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US-09-250-250E-26
US-09-250E-26
US-09-250E-26
Patent No. 683083
Patent No. 683084
Patent No. 683084
Patent No. 683084
CURRENT PILICATION NUMBER: US/09/269,250E
CURRENT PILICATION NUMBER: US/09/269,250E
NUMBER OF SEC ID NOS: 38
SOFTWARE: PatentIn version 3.1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               RESULT 6
US-09-269-250E-18
Sequence 18, Application US/09269250E
Sequence 18, Application US/09269250E
Sequence 18, Application US/09269250E
Sequence 18, Application US/09269250E
Patent No. 6830883
GENERAL INFORMATION:
HIGH APPLICANT GOUIN, ELSPA
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
TITLE OF INVENTION WINBER: US/09/269, 250E
CURRENT PELLING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
SEQ ID NO 18
LENGTH: 9
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                 TITLE OF INVENTION: Protein-Protein Complexes and Method of Using Same
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             OTHER INFORMATION: Description of Artificial Sequence: Exon Fragments
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                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 1148;
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                                                                                                                                                                                                                                                                                                                                                         LOCATION: (0)...(0)
COTHER INFORMATION: Polypeptide Accession Number YDR128W US-09-538-092-156
                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 38; DB 4;
Pred. No. 75;
                                FILE REPERINCE: 15966-542
CURRENT APPLICATION NUMBER: US/09/538,092
CURRENT FILING DATE: 2000-03-29
PURDA APPLICATION NUMBER: 60/127,352
PRIOR FILING DATE: 1999-04-01
PRIOR FILING DATE: 2000-02-01
NUMBER OF SEQ ID NOS: 1387
SOFTWARE: CURAPATSE CONDITION OF SEQ ID NOS: 1387
LENGTH: 1148
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      3; Mismatches
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66.7%;
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ORGANISM: Artificial Sequence
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1001 LLHDDIIEA 1009
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NAME/KEY: misc_feature
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Matches 8; Conserv
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LENGTH: 13
TYPE: PRT
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Requence 7115, Application US/09543681A

Patent No. 660709

RAPPLICANT: GARY BRETON

TITLE OF INVENTION: NUCLEIC ACID AND AMINO ACID SEQUENCES RELATING TO PROTEUS MIRABIL

TITLE OF INVENTION: DIAGNOSTICS AND THERAPEUTICS

FILE REFERENCE: 2709.1002-001

CURRENT APPLICATION NUMBER: US/09/543,681A

CURRENT APPLICATION NUMBER: US 60/128,706

PRIOR FILING DATE: 1999-04-09

NUMBER OF SEQ ID NOS: 8344

SEQ ID NO 3: 8344
                                                                                                                                                                                 RESULT 3
US-09-269-250E-36
Sequence 36, Application US/09269250E
Sequence 36, Application US/09269250E
Patent No. 6830883
GENERAL INFORMATION:
TITLE OF INVENTION:
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
TITLE REFERENCE: 58994
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
LENGTH: 26
LENGTH: 26
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                100.0%; Score 45; DB 4; Length 26; 100.0%; Pred. No. 0.064; tive 0; Mismatches 0; Indels
Pred. No. 0.031;
                                       0; Mismatches
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Patent No. 6753314;
GENERAL INFORMATION:
APPLICANT: Glot, Loic
APPLICANT: Mansfield, Traci A.
                 100.0%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CRGANISM: Proteus mirabilis US-09-543-681A-7115
                 Best Local Similarity 100.
Matches 9; Conservative
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16 VLHDDLLEA 24
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Matches 9; Conserv
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US-09-538-092-156
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GenCore version 5.1.6
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3, 2005, 06:53:47 ; Search time 17 Seconds June Run on:

(without alignments) 50.938 Million cell updates/sec

US-10-623-176-10

Perfect score:

1 VLRDDLLEA 9 Sequence:

Gapop 10.0 , Gapext 0.5 **BLOSUM62** Scoring table:

283416 seqs, 96216763 residues Searched:

283416 Total number of hits satisfying chosen parameters:

seq length: 0 seq length: 200000000 Minimum DB Maximum DB

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

PIR 79:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:* Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

	Description	C. elegans protein	protein K04A8.9 [i	type I restriction	transcription regu	probable membrane	glyceraldehyde-3-p	hypothetical prote	replicative DNA he	polyketide synthas	ABC transporter, m	hypothetical prote	surface adhesin A	quinone oxidoreduc	probable transposo	epidermal growth f	probable iles prot	isoleucyl-tRNA syn	two-component resp	hypothetical prote	nucleoid-asociated	hypothetical 37.8	nucleoid-associate	· nucleoid-associate	probable oxidoredu	iron-sulfur cofact	hypothetical prote	hypothetical prote		¤
SUMMARIES		33	75	92	11	28	43	04	78	48	18	58	14	72	73	28	20	28	32	78	96	98	57	13	53	99	11	30	69	37
ß		594	C89075	C69226	AI2271	831258	A64443	T20604	A82878	S43048	AG3118	H98168	D83714	T11672	H85073	138728	E70760	E87058	D70032	T41578	AC0786	A64988	H85857	F91013	D70753	T34766	AF2801	H97580	T40769	AD13
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	å Query Match Length	1165	174	616	192	282	343	389	486	3519	287	288	300	324	483	822	1041	1059	237	264	335	335	335	335	372	389	432	432	452	481
d	Query Match	100.0	85.7	83.3		81.0	81.0	81.0	81.0	81.0	78.6	78.6	78.6	78.6	78.6	78.6	78.6	78.6	76.2	76.2	76.2				ŝ	76.2	76.2	76.2	76.2	76.2
	Score	42	36	35	34	34	34	34	34	34	33	33	33	33	33	33	33	33	32	32	32	32	32	32	32	32	32	32	32	32
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leucyl aminopeptid	leucine aminopepti	GGDEF family prote	hypothetical prote	probable phosphori	hypothetical prote	hypothetical prote	. sulfite reductase	hypothetical prote	hypothetical prote	nitrate reductase					
G86538	G72083	AF2950	E98332	T00485	D83182	E71020	A34231	H65057	AG0858	C91081	D85926	AF0409	T46060	T46061	T02240
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499	499	525	525	545	268	587	599	599	599	599	599	909	826	826	883
76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2	76.2
32	32	32	32	32	32	32	32	32	32	32	32	32	32	35	32
30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

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C;Species: Homo sapiens (man)
C;Date: 03-Jun-2002 #sequence_revision 03-Jun-2002 #text_change 09-Jul-2004
C;Date: 03-Jun-2002 #sequence_revision 03-Jun-2002 #text_change 09-Jul-2004
C;Accession D594333
E;Nagase, T: Seki, N: Ishikawa, K.; Ohira, M.; Kawarabayasi, Y.; Ohara, O.; Tanaka, A.;
DNA Res. 3, 321-329, 1996
A;Title: Prediction of the coding sequences of unidentified human genes. VI. The coding s
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             A;Status: preliminary
A;Molecule type: DNA
A;Residues: 1-1165 «STO2>
A;Cross-references: GB:BAA13212; PID:g1504026; PIDN:BAA13212.1
                                                                                                                                                                                                                                       A; Reference number: D59433; MUID:97191544; PMID:9039502
elegans protein Z37093 homolog [imported] - human
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166 VLRDDLLEA 174
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protein K04A8.9 [imported] - Caenorhabditis elegans C;Species: Caenorhabditis elegans C;Date: 10-May-2001 #sequence_revision 10-May-2001 #text_change 09-Jul-2004

C; Accession: C89075
R; anonymous, The C. elegans Sequencing Consortium.
Schence 282, 2012-2018, 1998
A; Title: Genome Sequence of the nematode C. elegans: a platform for investigating biology A; Title: Genome Sequence of the nematode C. elegans: a platform for investigating biology A; Title: Genome Sequence of the nematode C. elegans: a platform for investigating biology A; Totle: See websites genome wustl.edu/gsc/C_elegans/ and www sanger.ac.uk/Projects/C_elegans/ A;Note: see websites genome wustl.edu/gsc/C_elegans/ and www sanger.ac.uk/Projects/C_elegans/ A;Accession: C89075
A;Accession: C89075
A;Accession: C89075
A;Accession: C8075
A;Residues: 1-174 <STO>
A;Residues: 1-174 <STO>
C;Genetics:

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probable membrane protein FUN34 - yeast (Saccharomyces cerevisiae)

NyAlternate names: protein N2029; protein YNR002C
C;Species: Saccharomyces cerevisiae
C;Date: 30-Sep-1993 #sequence_revision 30-Sep-1993 #text_change 09-Jul-2004
C;Accession: $31289, $45126; $48340, $63328
C;Accession: $31289, $45126; $48340, $63328
R;Stettler, S.; Mariotte, S.; Riva, M.; Sentenac, A.; Thuriaux, P.
J. Biol. Chem. 267, 21390-21395, 1992
A;Title: An essential and specific subunit of RNA polymerase III (C) is encoded by gene F A;Recession: $31288
A;Accession: $31288
A;Status: nucleic acid sequence not shown; translation not shown
A;Residues: 1-282 <STE>
A;Cross-references: UNIPROT:P32907; EMBL:X63746; NID:g4378; PIDN:CAA45279.1; PID:g4380
A;Note: the nucleotide sequence was submitted to the EMBL Data Library, February 1992
R;Verhasselt, P.; Aert, R.; Voet, M.; Volckaert, G.
submitted to the EMBL Data Library, January 1994
A;Pecenion: Twelve open reading frames revealed on the 23.6 kbp segment flanking the c
A;Accession: cariac
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C;Species: Methanococcus jannaschii
C;Date: 13-Sep-1996 #sequence_revision 13-Sep-1996 #text_change 09-Jul-2004
C;Accession: A6444;
R;Bult, C.J.; White, O.; Olsen, G.J.; Zhou, L.; Fleischmann, R.D.; Sutton, G.G.; Blake, C; Reich, C.I.; Overbeek, R.; Kirkness, E.F.; Weinstock, K.G.; Merrick, J.M.; Glodek, A.; rson, J.D.; Sadow, P.W.; Hanna, M.C.; Cotton, M.D.; Roberts, K.M.; Hurst, M.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   A,Accession: S45126
A,Molecule type: DNA
A,Rosidues: 1-282 <VERS-
A,Facilues: 1-282 <VERS-
A,Cross-references: EMBL:X77395; NID:g496717; PIDN:CAA54571.1; PID:g496720
R;Verhasselt, P.; Aert, R.; Voet, M.; Volckaert, G.
Yeast 10, 1355-1361, 1994
A,Title: Twelve open reading frames revealed in the 23.6 kb segment flanking the centrome A,Reference number: S48338; MUID:95208356; PMID:7900425
A,Accession: S48340
A;Status: nucleic acid sequence not shown; translation not shown
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A;Cross-references: EMBL:Z71617; NID:g1302470; PIDN:CAA96278.1; PID:g1302471; MIPS:YNR007
A;Experimental source: strain S288C
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A;Residues: 1-282 <VE2>
A;Residues: 1-282 <VE2>
A;Residues: 1-282 <VE2>
A;Cross-references: EMBL:X71395; NID:9496717; PIDN:CAA54571.1; PID:9496720
A;Note: the nucleotide sequence was submitted to the EMBL Data Library, January 1994
R;Aert, R; Verhasselt, P.; Voet, M.; Volckaert, G.
submitted to the Protein Sequence Database, April 1996
A;Reference number: S62910
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C;Superfamily: Saccharomyces cerevisiae probable membrane protein FUN34
C;Keywords: transmembrane protein
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Pred. No. 41;
0; Mismatches 1; Indels
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Best Local Similarity 87.5%;
Matches 7; Conservative
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R; Kaneko, T.; Nakamura, Y.; Wolk, C.P.; Kuritz, T.; Sasamoto, S.; Watanabe, A.; Iriguchi Nakazaki, N.; Shimpo, S.; Sugimoto, M.; Takazawa, M.; Yamada, M.; Yasuda, M.; Tabata, S DNA Res. 8, 205-213, 2001
A; Title: Complete Genomic Sequence of the Filamentous Nitrogen-fixing Cyanobacterium Ana A; Reference number: AB1807; MUID:21595285; PMID:11759840
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A;Residues: 1-192 «KUR»
A;Cross-references: UNIPROT:Q8YQT9; GB:BA000019; PIDN:BAB75427.1; PID:g17132862; GSPDB:G
A;Experimental source: strain PCC 7120
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            C, Accession: C69226
R; Smith, D.R.; Doucette-Stamm, L.A.; Deloughery, C.; Lee, H.; Dubois, J.; Aldredge, T.; Oliu, D.; Spadafora, R.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Gibson, R.; Jiwani, N.; Vicaire, R.; Wang, Y.; Wierzbowski, J.; Gibson, R.; Jiwani, N.; Vi. S.; Church, G.M.; Daniels, C.J.; Mao, J.; Rice, P.; Noelling, J.; Reeve, J.N.
J. Bacteriol. 179, 7135-7155, 1997
A; Title: Complete genome sequence of Methanobacterium thermoautotrophicum Delta H: funct A; Reference number: A69000; MUID:98037514; PMID:9371463
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    restriction modification enzyme, subunit M - Methanobacterium thermoautotrophicum
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A;Cross-references: UNIPROT:O27025; GB:AE000868; GB:AE000666; NID:g2622025; PIDN:AAB8544
A;Experimental source: strain Delta H
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A;Note: Nostoc sp. strain PCC 7120 is a synonym of Anabaena sp. strain PCC 7120
C;Date: 14-Dec-2001 #sequence_revision 14-Dec-2001 #text_change 09-Jul-2004
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C;Date: 05-Dec-1997 #sequence_revision 05-Dec-1997 #text_change 09-Jul-2004
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C;Superfamily: type I site-specific deoxyribonuclease chain hsdM
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.larity 100.0%; Pred. No. 27;
Conservative 0; Mismatches (
                                                                                                                       Score 36; DB 2;
Pred. No. 9.9;
2; Mismatches (
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Pred. No. 61;
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77.8%;
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ilarity 77.8%;
Conservative 1
                                                                                       Query Match
Best Local Similarity 77.8
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VLEDDLIEA 473
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129 VLKDDLLDA 137
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Matches 7; Conserv
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                                             A; Map position: 5
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A,Gene: all3728
A; Gene: K04A8.9
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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                                                                                                                                                                                                                                                                                                                                                                           1612378 segs, 512079187 residues
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Maximum Match 100%
Listing first 45 summaries
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## 133	00000000000000000000000000000000000000	RELIMINARY; PRT; 1131 AA. TEMBLE-1. 23, Created) THEMBLE-1. 25, Last sequence update) THEMBLE-1. 26, Last annotation update) PL1-associated RhoGAP 1 (Fragment). Human. Tazoa; Chordata; Craniata; Vertebrata; But anata; Primates; Catarrhini; Hominidae; Hob f; N.A. 187C. 187C. 19 P:intracellular signaling cascade; IRA. 200195 CytCals Fees CIP4. 200196; CACLS Fees CIP4. 200198; RhoGAP. 21. 1. 22. 1. 1. 23. RhoGAP; 1. 24. RhoGAP; 1. 25. FCH; 1. 26. CYTOCHROME C; UNKNOWN 1. 27. 1. 28. 1. 28. RhoGAP; 1. 29. 1. 20. 1. 2	core 42; red. No. Mismatch	PRT; 1136 13, Created) 3, Last sequency
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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SMART; SM00055; FCH; 1.
SMART; SM00324; RhoGAP; 1.
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166 VLRDDLLEA 174
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                NCBI_TaxID=9606;
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                                                                                SEQUENCE FROM N.A.
MEDLINE=20549028; PubMed=11095984; DOI=10.1006/bbrc.2000.3880;
Kaminski W. B., Pichler A., Schmitz G.;
"Genomic organization of the human cholesterol-responsive ABC
transporter ABCA7: tandem linkage with the minor histocompatibility
antigen HA-1 gene.";
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01-FEB-1997 (TrEMBLrel. 02, Last sequence update)
01-FBB-1997 (TrEMBLrel. 26, Last annotation update)
KIAA0223 protein (Fragment).
Name=KIAA0223, Human).
Homo sapiens (Human).
Eukaryota, Metazoa, Chordata, Craniata, Vertebrata; Euteleostomi;
                 Eukaryota, Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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InterPro; IPR001066; GG15. Fee CIP4.
InterPro; IPR000345; CytC heme BS.
InterPro; IPR002219; DAG_PE-bind.
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EMBL; AF308045; AAN04658.1; JOINED.
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PROSITE; PS00479; DAG PE BIND DOM 1; UNKNOWN 1.
PROSITE; PS500181; DAG PE BIND DOM 2; 1.
SEQUENCE: PS50738; RHOĞAP; 1.
SEQUENCE: 1136 AA; 124613 MW; 9536787B3B1EE1
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Interpro; IPR008936; Rho_GAP
Pfam; PF00130; Cl_1; 1.
Pfam; PF00620; RhoGAP; 1.
                                                                                                                                                                                                                            AF308046; AAN04658.1;
                                                                                                                                                                                                                                                            AF308048; AAN04658.1;
AF308049; AAN04658.1;
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AAN04658.1;
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SMART; SM00055; FCH; 1.
SMART; SM00324; RhoGAP; 1.
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 Homo sapiens (Human)
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Best Local Similarity
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                                                  NCBI_TaxID=9606;
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TISSUE=Bone marrow;

X MEDLINE=S7191944; Pubmed=9039502;

A Radiase T. Seki N., Ishikawa K., Ohira M., Kawarabayasi Y., Ohara O.,

A Tanaka A., Kotani H., Miyajima N., Nomura N.;

Tanaka A., Kotani H., Miyajima N., Nomura N.;

Tanaka A., Kotani H., Miyajima N., Nomura N.;

The coding sequences of unidentified human genes. VI.

The coding sequences of 80 new genes (KIAA0201-KIAA0280) deduced by

Tanalysis of CDNA clones from cell line KG-1 and brain.";

In DNA Res. 3:321-329(1996).

R PRB: D59433; D59433.

R PRB: D59433; D59433.

R PSR: PSR: Pintracellular signaling cascade; IEA.

R Pfam: PF00130; Cl 1:1.

R Pfam: PF00130; Cl 1:1.

Pfam: PF00120; KhoGAP; 1.
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Parasitiformes; Ixodida; Ixodidae; Ixodes.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    STRAIN=Rhode Island; TISSUE=Salivary gland; Valenzuela J.G., Francischetti I.M., Pham V.M., Garfield M., Mather T.N., Ribeiro J.M.C.; Submitted (FRB-2002) to the EMBL/GenBank/DDBJ databases.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            100.0%; Score 42; DB 2; Length 1165; 100.0%; Pred. No. 31; 2; ve 0; Mismatches 0; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Hydrolase, Protease, Serine protease. –
SEQUENCE 376 AA, 41434 MW, 64F403CAD505CD33 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               01-07T-2002 (TrEMBLrel. 22, Created)
01-0CT-2002 (TrEMBLrel. 22, Last sequence update)
01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Putative serine protease with signal anchor.
Ixodes scapularis (Black-legged tick) (Deer tick)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  PROSITE; PS00190; CYTOCHROME C; UNKNOWN 1.
PROSITE; PS00419; DAG PE BIND DOM 1; UNKNOWN 1.
PROSITE; PS50081; NHGGAP; 11.
PROSITE; PS50238; RHGGAP; 11.
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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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June Run on:

3, 2005, 06:44:47; Search time 90.5 Seconds (without alignments) 38.462 Million cell updates/sec

US-10-623-176-10

1 VLRDDLLEA 9 Perfect score: Sequence: BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

2105692 segs, 386760381 residues Searched:

2105692 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 200000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

A_Geneseq_16Dec04:* 1: geneseco1980r.* .. Database

geneseqp1990s:* geneseqp2000s:* geneseqp2001s:* geneseqp20028:* geneseqp2004s:*

geneseqp2003as:* geneseqp2003bs:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		æ			SOMMEN		
Result No.	Score	Query	Length	DB	ID	Description	
7	42	100.0	6	2	AAW99197	Aaw99197 Minor his	
8	42	100.0	6	7	AAW97374	Aaw97374 HA-1 R-al	
m	42	100.0	6	œ	ADH40334	Adh40334 Human min	
4	42	100.0	13	~	AAW99198	Aaw99198 DH cell K	
ហ	42	100.0	13	~	AAW97414	Aaw97414 KIAA0223	
9	42	100.0	1136	ß	ADI17193	Adil7193 Human NOV	
7	42	100.0	1136	ß	ADI16744	Adil6744 Human NOV	
80	42	100.0	1136	80	ADH61299	Adh61299 INTSIG pr	
6	42	100.0	1136	œ	ADN42398	Adn42398 Human nov	
10	42	100.0	1136	8	ADS10490	Ads10490 Human the	
11	42	100.0	1165	Ŋ	ADI17194	Adil7194 Human NOV	
12	42	100.0	1165	œ	ADP55035	Adp55035 Human PRO	
13	42	100.0	1165	œ	ADP24600	Adp24600 PRO polyp	
14	38	90.5	4150	m	AAY92707		
15	37	88.1	6	~	AAW99196	Aaw99196 Minor his	
16	37	88.1	6	N	AAW97375	Aaw97375 HA-1 H-al	
17	37	88.1	6	æ	ADH40333	Adh40333 Human min	
18	37	88.1	13	~	AAW99199	Aaw99199 vR cell K	
19	37	88.1	13	7	AAW97415	Aaw97415 KIAA0223	
20	37	88.1	1165	ω	ADS11768	Ads11768 Human the	
21	37	88.1	1177	ø	ABP71254	Abp71254 Chimeric	
22	37	88.1	1186	9	ABP71253	Abp71253 Zebrafish	
23	36	85.7	6	7	AAW99195	_	
24	36	85.7	σ	~	AAW97572	٠.	
25	34	81.0	282	S	ABG93191	Abg93191 S. cerevi	

Aaus1225 Propionib Abm47744 Propionib Adni8333 Bacterial				Abu34693 Protein e Abu36647 Protein e Aaw57448 M. tuberc Abu35902 Protein e
AAU51225 ABM47744 ADN18393	AAY9270 AAY92709 AAY92708 AAU35037	AAG73861 ADC07956 ADC08309	ADCUGES ADS29025 ABP58031 ADN95677 AAE06696	ABU3 6647 ABU3 6647 AAW 57448 ABU3 5902
324 324 343 6		328 4 472 7 486 7	491 738 822 6822 6822 7	041 6 041 6 045 2 059 6
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ALIGNMENTS

Minor histocompatibility antigen HA-1 T-cell epitope #3 AAW99197 standard; peptide; 9 AA. (first entry) 20-MAY-1999 AAW99197; RESULT 1 AAW99197

Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.

Homo sapiens

WO9905174-A1

04-FEB-1999.

98WO-NL000425. 23-JUL-1998; 97EP-00202303. 23-JUL-1997;

(UYLE-) RIJKSUNIV LEIDEN

Engelhard VH;

Hunt DF,

Goulmy EAJM,

WPI; 1999-153312/13.

A new minor histocompatibility antigen, HA-1 - useful to treat immune diseases and prevent rejection and host versus graft disease in bone marrow and organ transplantation.

Disclosure; Page 15; 47pp; English.

The present sequence represents a new peptide (P1) constituting a T-cell epitope obtainable from the minor histocompatibility antigen HA-1. The peptide is immunogenic and can be used as part of a vaccine. P1 is used as a medicine, to induce tolerance for transplants, prevent rejection and/or graft versus host disease, or to treat (auto) immune diseases. In particular it can be used with bone marrow transplantation, in the treatment of severe aplastic anaemia, leukaemia, and immune deficiency

Sequence 9 AA;

Query Match

100.0%; Score 42; DB 2; Length 9;

RESULT 2 AAW97374

8

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The invention relates to a novel method for providing epitopes of allelic variants of antigenic proteins from specific species based on single nucleotide polymorphism (SNP), by defining target protein/jeptide or its subset, screening database of DNA encoding target protein, identifying, selecting allelic peptide/protein variants, expression product or its fragment encoded by DNA sequence having SNP, creating variant epitopes, confiding to MHC protein. A protein of the invention has cytostatic activity, and may have a use in a vaccine. The method is useful for generating a SNP profile of one or more individuals from a given species by applying the method for several protein from the individuals, where the SNP profile was related to disease, preferably cancer. This is useful for diagnosing a disease in an individual by cancer. This is useful for diagnosing a disease in an individual by cancer. This is useful for tanaplanting haematopoietic stem cells from a donor to a recipient and treating cancer, preferably leukaemia, and for determining the progression, regression conset of a treated disease. The present consequence is used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Providing allelic variant epitope of protein based on single nucleotide polymorphism by defining target protein, screening database of protein, identifying, selecting allelic variant protein, creating variant
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.
                                                                                     human, cytostatic, vaccine, SNP profile, cancer, leukaemia, minor histocomptibility antigen, mHAg; T cell epitope.
                                         Human minor histocomptibility antigen HA-1 T cell eptiope
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Disclosure; Page 82; 119pp; English.
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                                                                                                                                                                                                                                                                                                                                      13-JUN-2002; 2002EP-00013423.
                                                                                                                                                                                                                                                                                          3-JUN-2003; 2003WO-EP006251.
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  11-MAR-2004 (first entry)
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Les 9; Conservative
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                                                                                                                                                            Homo sapiens.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 The present sequence represents part of the minor histocompatibility antigen HA-1 R-allele. The specification describes methods for typing alleles (preferably the H and R alleles) of the minor histocompatibility antigen HA-1 in a sample, which comprise detecting polymorphic nucleotides in the cDNA or genomic nucleic acids of the alleles. The methods can be used for HA-1 typing for bone marrow transplants, severe aplastic anaemia, leukaemia and immune deficiency diseases, as well as detection of genetic aberrances. The probes and primers of the invention can be used to screen for the HA-1 alleles. The HA-1 peptides can be used anti-idiotypic B cells and/or T cells and antibodies
                                                                                                                                                                                                                                                                                                                                                                                                    Intron, minor histocompatibility antigen HA-1; typing allele; H allele, R allele; polymorphic nucleotide; HA-1 typing; bone marrow transplant; severe aplastic anaemia; leukaemia; immune deficiency disease; ss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Typing minor histocompatibility antigen HA-1 - by amplifying and identifying HA-1 R and HA-1 H alleles of HA-1, used for, e.g. detection
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                       Indels
    Pred. No. 1.8e+06;
                       0; Mismatches
                                                                                                                                                                                                                          AAW97374 standard; protein; 9 AA.
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100.08;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               98WO-EP004928
                                                                                                                                                                                                                                                                                                                  13-MAY-1999 (first entry)
                         9; Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         (UYLE-) RIJKSUNIV LEIDEN
                                                                                                                                                                                                                                                                                                                                                             HA-1 R-allele sequence
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    Best Local Similarity
Matches 9; Conserv
                                                                  1 VLRDDLLEA
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          WO9905313-A2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               23-JUL-1998;
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02-JUN-1998;
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                                                                                                                                                                                                                                                                      AAW97374;
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Gaps ö

ADH40334 standard; peptide; 9 AA.

RESULT 3

ADH40334

Best Loc Matches

ADH40334;

Length 9; 0; Indels

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GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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June Run on:

3, 2005, 07:09:04; Search time 70 Seconds (without alignments) 44.445 Million cell updates/sec

US-10-623-176-10 42

1 VLRDDLLEA 9 Title: Perfect score: Seguence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

1465611 segs, 345679903 residues Searched:

1465611 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

Published Applications AA:*

1: \cgn2 \(\) \cgn2 \(\) \(\) \(\) \cgn2 \(\) \(\) \cgn2 \(\) \(\) \(\) \cgn2 \(\) \cgn2 \(\) \cgn2 \(\) \cgn2 \(\) \cgn2 \(\) \(\) \(\) \cgn2 \(\) \(\

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

	Appl	Appli	Appl										
_	10,	5,	44,	65,	74,	14,	69	70,	71,	63,	72,	67,	99
Description	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Seguence	Sequence	Sequence	Sequence	Sequence	Sequence
Œ	US-10-623-176-10	US-10-791-217-5	US-10-623-176-44	US-10-623-176-65	US-10-623-176-74	US-10-791-217-14	US-10-623-176-69	US-10-623-176-70	US-10-623-176-71	US-10-623-176-63	US-10-623-176-72	US-10-623-176-67	US-10-623-176-66
	15	16	15	15	15	16	15	15	15	15	15	15	15
* Query Match Length DB	6	Đ	10	12	13	13	17	19	21	22	23	25	56
& Query Match	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0	100.0
Score	42	42	42	42	42	42	42	42	42	42	42	42	42
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Sequence 60, Appl	61,	29	28	72	73	42	46	'n	ď	ď	'n		16,	16,	57,	23	22	51,	49	4,	ď	ų	'n	Sequence 24, Appl	33	48	34	유			Sequence 76901, A
US-10-623-176-60	US-10-623-176-61	-10-623-176-	-10-072-012	-10-012	-10-072-	US-10-623-176-42	- 1	US-09-808-880-2	US-10-623-176-2	US-10-791-217-2	US-10-861-335-1	-10-623-	0-623	0 - 791 -	-10-623-	US-10-623-176-53	0-623-	0-623-176-5	3-623-	US-10-193-692-4	US-10-193-692-2	-10-623	0-791-	0-623	-10-623-	US-10-623-176-48	US-10-451-467A-340	-10-369-493-1	-10-437-963	US-10-437-963-114620	US-10-282-122A-76901
15	15	15	15	15	15	15	15	10	15	16	11	15	15	16	15	15	15	15	15	14	14	15	16	15	15	15	16	15	16	16	12
27	28	53	1136	1136	1165	σ	σ	4150	σ	Q	σ	10	13	13	17	22			53	1177	1186	σ	σ	10	17	0	282	343	407	408	486
100.0	100.0	100.0	100.0	100.0	100.0	90.5	90.5	90.5	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	88.1	85.7	85.7	85.7	85.7	81.0	81.0	81.0	81.0	81.0	81.0
42	42	42	42	42	42	38	38	38	37	37	37	37	37	37	37	37	37	37	37	37	37	36	36	36	36	34	34	34	34	34	34
14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36	37	38	39	40	41	42	43	44	45

ALIGNMENTS

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FEATURE:
OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
Sequence 10, Application US/10623176

Sequence 10, Application US/10623176

Publication No. US20040092446A1

GENERAL INFORMATION:
APPLICANT: Goulmy, Els A.J.M.
APPLICANT: Hunt, Donald F.

TITLE OF INVENTION: HA-1 epitopes and uses thereof
FILE REFERENCE: 2183-6047US

CURRENT APPLICATION NUMBER: US/10/623,176

CURRENT FILING DATE: 2003-07-18

PRIOR FILING DATE: 2000-01-21

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-24

PRIOR FILING DATE: 1998-07-24

NUMBER OF SEQ ID NOS: 77

SEQ ID NOS: 77

LENGTH: 9

LENGTH: 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TYPE: PRT
ORGANISM: Artificial Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ; NAME/KEY: SITE
; LOCATION: (1)..(9)
US-10-623-176-10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      FEATURE:
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Gaps ö Query Match
100.0%; Score 42; DB 15; Length 9;
Best Local Similarity 100.0%; Pred. No. 1.3e+06;
Matches 9; Conservative 0; Mismatches 0; Indels

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US-10-623-176-65
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OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ; OTHER INFORMATION: obtained from KIAA0223 partial complementary DNA
US-10-791-217-5
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Sequence 44, Application US/10623176

; Publication No. US20040092446A1

; GENERAL INFORMATION:

APPLICANT: Goulmy, Els A.J.M.

; APPLICANT: BENGENAL R.

; TITLE OF INVENTION: HA-1 epitopes and uses thereof

; TITLE OF INVENTION: HA-1 epitopes and uses thereof

; TITLE OF INVENTION: HA-1 epitopes and uses thereof

; FILE REFERENCE: 2183-6047US

; CURRENT APPLICATION NUMBER: US/10/623,176

; CURRENT FILING DATE: 2003-07-18

; PRIOR APPLICATION NUMBER: EP 97202303.0

; PRIOR APPLICATION NUMBER: EP 97202303.0

; PRIOR PILING DATE: 1999-07-23

; PRIOR FILING DATE: 1998-07-23

; PRIOR FILING DATE: 1998-07-23

; PRIOR FILING DATE: 1998-07-23

; PRIOR FILING DATE: 2000-01-24

; SOFTWARE: OF SEQ ID NOS: 77

; SOFTWARE: PRIOR 140
                                                                                                                                                                                                                            CEMERAL INFORMATION:
APPLICANT: Goulmy, Elsa A.J.M
APPLICANT: Hunt, Donald F
APPLICANT: Hunt, Donald F
APPLICANT: Hunt, Donald F
TILE REFERENCE: 2183-4285US
CURRENT APPLICATION NUMBER: US/10/791,217
CURRENT APPLICATION NUMBER: US/09/489,760
PRIOR FILING DATE: 2004-03-02
PRIOR FILING DATE: 2000-01-21
PRIOR PILING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: ParentIn Ver. 2.1
SENGTH: 9
                                                                                                                                                              Sequence 5, Application US/10791217
Publication No. US20040191268A1
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               TYPE: PRT
ORGANISM: Artificial sequence
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1 VLRDDLLEA
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NAME/KEY: SITE
LOCATION: (1)..(10)
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100.0%; Score 42; DB 15; Length 10; 100.0%; Pred. No. 0.2;
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                                                   0; Indels
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Sequence 74, Application US/10623176

Publication No. US20040092446A1

GENERAL INFORMATION:
APPLICANT: Goulmy, Els A.J.M.
APPLICANT: Hunt, Donald F.
TITLE OF INVENTION: HA-1 epitopes and uses thereof
FILE REFERENCE: 2183-6047US

CURRENT APPLICATION UNMER: US/10/623,176

CURRENT FILING DATE: 2003-00-18

PRIOR FILING DATE: 2000-01-21

PRIOR FILING DATE: 1997-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR APPLICATION NUMBER: PCT/NL98/00424

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1900-01-24

NUMBER OF SEQ ID NOS: 77
                                                                                                                                                                                                                                                                                                                                                                GENERAL INFORMATION:
APPLICANT: GOULMY, E18 A.J.M.
APPLICANT: Hunt, Donald F.
APPLICANT: Hunt, Donald F.
TITLE OF INVENTION: Ha-1 epitopes and uses thereof;
FILE REPRENCE: 2183-6047US
CURRENT APPLICATION NUMBER: US/10/623,176
CURRENT PILING DATE: 2003-07-18
PRIOR APPLICATION NUMBER: 09/489,760
PRIOR APPLICATION NUMBER: EP 97202303.0
PRIOR FILING DATE: 1997-07-23
PRIOR FILING DATE: 1988-07-23
PRIOR FILING DATE: 1988-07-23
PRIOR FILING DATE: 1997-07-23
PRIOR FILING DATE: 2000-01-24
NUMBER OF SEQ ID NOS: 77
SOFTWARE: PATENTING DATE: 2000-504165
PRIOR FILING DATE: 2000-01-24
NUMBER OF SEQ ID NOS: 77
SOFTWARE: PATENTING DATE: 2000-61-24
SEQ ID NO 65
LENGTH: 12
                                                                                                                                                                                                                                                                                           ; Sequence 65, Application US/10623176; Publication No. US20040092446Al; GENERAL INFORMATION:
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ORGANISM: Artificial Sequence
Query Match 100.
Best Local Similarity 100.
Matches 9; Conservative
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; LOCATION: (1)..(12)
US-10-623-176-65
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10, Appl 6729, Ap 22, Appl

Sequence Sequence Sequence Sequence

Sequence Sequence Sequence 22, Appl 20462, A 23, Appl 31013, A 42, Appl

Sequence

Sequence Sequence Sequence Sequence Sequence Sequence Sequence

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Run on:

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Searched:

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Sequence 26, Application US/09269250E
Sequence 26, Application US/09269250E
Sequence 26, Application US/09269250E
Sequence 26, Application US/09269250E
PAPLICANT: Goulmy, Elsa
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REFERENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269,250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
SEQ ID NO 26
LENGTH: 13
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TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REFERENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269,250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ 1D NOS: 38
SOFTWARE: PATENTIN VEFSION 3.1
SEQ 1D NO 18
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US-10-065-133A-5
US-10-434-811A-2
US-10-434-811A-2
US-09-540-236-3485
US-09-9102-540-12839
US-08-9102-540-12839
US-08-396-001-10
US-08-396-001-10
US-09-323-433A-10
US-09-323-433A-10
US-09-134-000C-6729
US-09-134-035-425
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100.0%; Score 42; DB 4; I
Best Local Similarity 100.0%; Pred. No. 4.1e+05;
Matches 9; Conservative 0; Mismatches 0;
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ORGANISM: Artificial Sequence
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ORGANISM: Artificial Sequence
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   US-09-269-250E-18
   LENGTH: 9
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Sequence 404, App
Sequence 33603, A
Sequence 30099, A
Sequence 10099, A
Sequence 10059, A
Sequence 10059, A
Sequence 1, Appli
Sequence 2, Appli
Sequence 5, Appli
Sequence 5, Appli
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27.993 Million cell updates/sec
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/cgn2_6/ptodata/1/iaa/5B_COMB.pep:*
/cgn2_6/ptodata/1/iaa/6A_COMB.pep:*
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/cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
/cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*
                 GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-506-286B-5
US-09-762-861B-2
US-09-762-861B-5
US-10-065-133A-2
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Maximum Match 100%
Listing first 45 summaries
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Maximum DB seq length: 200000000
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Match Length DB
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Matches
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                                                                                                                                                                                        Sequence 36, Application US/09269250E
Patent No. 6830883
GENERAL INFORMATION:
APPLICANT: GOLIMY, E18a
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
CURRENT APPLICATION WUMBER: US/09/269,250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
SEQ ID NO 36
LENGTH: 26
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Betent No. 6231636

GENERAL INFORMATION:
APPLICANT: Betlach, Mary C.
APPLICANT: Tang, Li
TITLE OF INVENTION: RECOMBINANT OLEANDOLIDE POLYKETIDE SYNTHASE
FILE REFERENCE: 30062-20029, 00
CURRENT APPLICATION NUMBER: US/09/428,517
CURRENT FILING DATE: 1999-10-28

BEALLIER APPLICATION NUMBER: 60/120,254

EARLIER PILING DATE: 1999-02-16

BARLIER PILING DATE: 1999-02-16

BARLIER FILING DATE: 1999-02-16

BARLIER FILING DATE: 1998-10-29

NUMBER OF SEQ ID NO.S: 12
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US-09-269-250E-36
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US-09-428-517-2
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                    0; Indels
  Pred. No. 0.025;
                        0; Mismatches
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100.08;
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ORGANISM: Artificial Sequence
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SEQ ID NO 2
LENGTH: 4150
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Best Local Similarity 100.
Matches 9; Conservative
                      9; Conservative
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Best Local Similarity
Matches 8; Conserv
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US-09-269-250E-36
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Sequence 29, Application US/09269250E
Sequence 29, Application US/09269250E
Sequence 29, Application US/09269250E
GENERAL INFORMATION:
APPLICANTY: GOLIMY, ELSA
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1;
FILE REFERENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269,250E
CURRENT PILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: PALENTIN version 3.1
SEQ ID NO 29
LENGTH: 9
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US-09-269-250E-28
Squence 28, Application US/09269250E
Sgewence 28, Application US/09269250E
Spatent No. 6830883
GENERAL INFORMATION:
APPLICANT GOULD,
STILE OF INVENTION:
TILLE OF INVENTION:
TILLE OF INVENTION:
CURRENT APPLICATION NUMBER: US/09/269,250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: PatentIn version 3.1
                                                                                                        GENERAL INFORMATION:

BAPLICANT: GOUINY, BLSa

TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1

FILE REPERENCE: 58994

CURRENT APPLICATION NUMBER: US/09/269, 250E

CURRENT FILING DATE: 1999-05-21

NUMBER OF SEQ ID NOS: 38

SOFTWARE: Patentin version 3.1

SEQ ID NO 20

LENGTH: 9
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RESULT 5
US-09-266-250E-20
F Septence 20, Application US/09269250E
Patent No. 6830883
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ORGANISM: Artificial Sequence
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Matches 8; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              791
GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    283416 segs, 96216763 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            SUMMARIES
                                                                                                                                       OM protein - protein search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Listing first 45 summaries
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      S55696
P00663
P00663
S78764
S25948
S25948
A12876
A1
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PT0722
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PT0368
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Post-processing: Minimum Match 0% Maximum Match 100%
                                                                                                                                                                                                                                                                                                                                                     US-10-623-176-2
                                                                                                                                                                                                                                                                                                                                                                                                                1 VLHDDLLEA 9
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1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Minimum DB seq length: 0 Maximum DB seq length: 9
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 BLOSUM62
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No.
                                                                                                                                                                                                                                                                                                                                                     Title:
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phosphoenolpyruvate carboxykinase - Trypanosoma brucei
C;Species: 28-0ct-1995 #sequence_revision 03-Nov-1995 #text_change 09-Jul-2004
C;Accession: S55696
R;Hunt, M.; Koehler, P.
Biochim. Biophys. Acta 1249, 15-22, 1995
A;Title: Purification and characterization of phosphoenolpyruvate carboxykinase from Tryk
A;Reference number: S55696; MUID:95284106; PMID:7766679
A;Accession: S55696
A;Accession: S55696
A;Accession: S55696
A;Accession: S5696; MUID:95284106; PMID:7766779
A;Accession: S5696; MUID:95284106; PMID:7766779
A;Accession: S5696; MUID:95284106; PMID:7766779
A;Accession: S5696; MUID:95284106; PMID:7766779

RESULT 2

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Gaps

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37.8%; Score 17; DB 2; Length 9; llarity 33.3%; Pred. No. 2.8e+05; Conservative 3; Mismatches 1; Indels

Query Match Best Local Similarity Matches 2; Conserv

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RESULT PQ0663

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	32	77	7.97	- 1	ס ע	N	000	B30572	T-cel.	rec	eptor be		
	55.	77	7.97		י ע	7	APO	Abolius	exoco	מון א	exocoxin A - strep		
	34	;	24.4	4	n ı	7	2 1	C41225	cobbe	i rea	copper resistance		
	35	11	24.4	4	v.	7	110	T10954	nypoti	jeric	hypotherical prote		
	36	11	24.4	4	9	N	111	T11779	phospi	y1got	phosphoglycerate t		
	37	11	24.4	4	7	7	\$25	S25266	pilE 1	prote	pilE protein - Esc		
	38	11	24.4	4	7	04	PT0	PT0246	Ig heavy chain	avy o	hain CRD		
	39	11	24.4	4	8	~	\$22	S22428	chitin-binding	rid-۲	ding pro		
	40	11	24.4	4	8	8	B33	B33099	158K	axoan	158K exoantigen -		
	41	11	24.4	4	80	~	869	\$69165	ferre	loxin	ferredoxin až - Ja		
	42	11	24.4	4	0	7	PTO	PT0272	Iq he	S V	Iq heavy chain CRD		
	43	11	24.4	4	σ	~	998	S66419	tetran	neric	tetrameric protein		
	44	11	24.4	4	σ	7	A11	A11497	trans	ldol	transaldolase (EC		
	45	11	24.4	4	Φ	7	B39	B39504	octame	ır-bi	octamer-binding pr		
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							•	ALIGNMENTS					
	RESULT 1												
	I46868												
	alpha-myosin heavy chain - rabbit	osin he	avy c	hain	- ra	bbi	T E	(fragment)					
	C;Specie	B: Oryc	tolag	ns cn	nico	lus	. ق	C; Species: Oryctolagus cuniculus (domestic rabbit)	•		,		
	C;Date: 14-Feb-1997	14-Feb-	1997	#sedn	ence	re	V181	C;Date: 14-Feb-1997 #Bequence_revision 14-Feb-1997 #text_change 09-Jul-2004	t_change (ט-ענ	T-2004		
	C;Access P.Friedm	בי ער הפי		o Po	b D	ė.	e du	H					
_	K; FITEUM	יים לונה) i	י ברשי		2	o C	K;FILECUMAII, D.O.; UMECA, F.N.; SILLIA, A.M.; DBU, D.					
	FIOC. NA	CL. ACA			. ų	d i	7 1	Frice. Natl. Acad. Sci. 0.3.A. ol., 3044-50460, 1304.	and the sale	4	nad horr	4400	;
_	A;IICIE:	Cliatac	7777	TAFBE	. a	בולות בולות בולות	. 842	Ajiicie: Chaiaccellzación of genomic clones specifying 1 A.Doforonco mumbor. 146869. MITD.04221001. DMTD.6328491	ומחחור מזן	1011	מוות חברם	- אפוורד דרמדם	;
	A.Acression: T46868	ion. TA	6868		•		:	14.001100.001111				_	
	A:Status	preli	ninar	V: tr	ansl	ate	dfr	A:Status: preliminary: translated from GB/EMBL/DDBJ					
	A: Molecule type: DNA	le type	: DNA										
	A:Residues: 1-7 <fri></fri>	es: 1-7	<fri< td=""><td>٨</td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td>•</td><td></td></fri<>	٨								•	
	A; Cross-	referen	ces:	UNIPR	Q: TO	287	42;	A;Cross-references: UNIPROT:Q28742; GB:K01698; NID:g165538; PIDN:AAA31415.1; PID:g165539	538; PIDN	: AAA3	1415.1;	PID:9165539	0
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	Query Match	Match			40.	40.04;			Length 7;				
	Best Local			larity 50.	50.	50.08;	c	ed. No. 2.8e+05; Mismatches	Tndela		aneg	·	
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	q	3 Ж	3 MHDE 6										

Best Loc Matches

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elastase - Pseudomonas aeruginosa
C;Species: Pseudomonas aeruginosa
C;Date: 19-Mar-1997 #sequence_revision 24-Jul-1997 #text_change 24-Jul-1997
C;Accession: S20446
R;Kessler, E.; Safrin, M.; Peretz, M.; Burstein, Y.
FEBS Lett. 299, 291-293, 1992
A;Title: Identification of cleavage sites involved in proteolytic processing of Pseudomor A;Reference number: S20446; MUID:92183956; PMID:1544509
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ö
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                                                                                                                                                                                                                                                                                                                                                                                               A;Molecule type: protein
A;Residues: 1-6 <GRA>
C;Keywords: mitochondrion
F;1-6/Product: ribosomal protein MRP-S23 (fragment) #status experimental <MAT>
                                                                                                                                          ribosomal protein MRP-S23, mitochondrial - bovine (fragment)
C;Species: Bos primigenius taurus (cattle)
C;Bate: 20-Sep-1999 #sequence_revision 20-Sep-1999 #text_change 20-Sep-1999
C;Accession: S78764
R;Graack, H.R.
submitted to the Protein Sequence Database, July 1999
A;Reference number: S78760
A;Accession: S78764
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      C;Species: Rattus norvegicus
C;Date: 25-Aug-2003 #sequence_revision 25-Aug-2003 #text_change 25-Aug-2003
C;Accession: A59489
R;Negoro, M.
R;Negoro, M.
Rbantited to the Protein Sequence Database, June 2003
A;Description: Purification of PKCI from rat liver.
A;Reference number: A59489
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query Match 33.3%; Score 15; DB 2; Length 7; Best Local Similarity 40.0%; Pred. No. 2.8e+05; Matches 2; Conservative 1; Mismatches 2; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Length 6;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            A;Status: preliminary
A;Molecule type: protein
A;Residues: 1-7 <NEG>
A;Experimental source: strain Wistar, liver
A;Note: p-Hydroxyacetophenone-Sepharose binding protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 33.3%; Score 15; DB 2; L Best Local Similarity 75.0%; Pred. No. 2.8e+05; Matches 3; Conservative 0; Mismatches 1;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          protein kinase C inhibitor - rat (fragment)
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A;Molecule type: protein
A;Residues: 1-7 <KES>
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         A; Accession: A59489
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              RESULT 8
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R;Edgar, P.F.
RsEdgar, P.F.
A;Title: Hucolin, a new corticosteroid-binding protein from human plasma with structural A;Reference number: S68004; MUID:96087107; PMID:7498469
membrane protein - porcine epidemic diarrhea virus (isolate Belgian CV777) (fragment)
C;Species: porcine epidemic diarrhea virus
C;Date: 14-Uul-1994 #sequence_revision 14-Uul-1994 #text_change 08-Oct-1999
C;Accession: PQ0663
R;Bridgen, A; Duarte, M.; Tobler, K.; Laude, H.; Ackermann, M.
J. Gen. Virol. 74, 1795-1804, 1993
A;Title: Sequence determination of the nucleocapsid protein gene of the porcine epidemic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    C;Accession: PC4131

K;Kawasaki, S.; Arai, H.; Igarashi, Y.; Kodama, T.

Gene 167, 87-91, 1995

A;Title: Sequencing and characterization of the downstream region of the genes encoding y for biosynthesis of heme d1.

A;Reference number: JC4552; MUID:96144254; PMID:8566817

A;Accession: PC4131
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C,Species: Homo sapiens (man)
C,Date: 15-Feb-1997 #sequence_revision 13-Mar-1997 #text_change 17-Mar-1999
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         hypothetical protein 8 [imported] - Pseudomonas aeruginosa (fragment)
C.Species: Pseudomonas aeruginosa
C.Date: 20-Apr-2000 #sequence_revision 20-Apr-2000 #text_change 09-Jul-2004
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        A;Cross-references: UNIPROT:P95412; DDBJ:D50473; NID:g1217594
A;Note: this ORF is not annotated in GenBank entry PSENIRC, release 113.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gaps
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                                                                                                                                                                                                                                                                                                                                                                   A;Cross-references: GB:214976; NID:g311650; PIDN:CAA78699.1; PID:g584083 C;Comment: This virus is coronavirus related to human coronavirus 229E; C;Keywords: membrane protein
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                                                                                                                                                                                                            issible gastroenteritis virus.
A;Reference number: JQ2191; MUID:93389433; PMID:8397280
A;Accession: PQ0663
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Best Local Similarity 100.
Matches 3; Conservative
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A;Molecule type: protein
A;Residues: 1-7 <EDG>
                                                                                                                                                                                                                                                                                                             A; Molecule type: mRNA A; Residues: 1-7 < BRI>
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GenCore version 5.1.6
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June 3, 2005, 07:11:54; Search time 77.5 Seconds (without alignments) 59.467 Million cell updates/sec Run on:

US-10-623-176-2 45 1 VLHDDLLEA 9

Title: Perfect score: Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

1612378 segs, 512079187 residues Searched:

1455 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 9

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

UniProt_03:*
1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		Description	Q7r8x5 plasmodium	Q47556 escherichia	Q28742 oryctolagus	Q7m3s5 trypanosoma	099182 gnatholebia	Q7z6g0 homo sapien	P83281 macrobrachi	Q70sm2 homo sapien	Q8lpt5 zea mays (m	strei	_	bacte	P36414 pseudomonas	P02729 homo sapien	Q9bfa7 macroscelid	P72279 rhodococcus	_	P17440 pichia jadi	homo	•		-	Q8cg39 rattus norv	Q9qza8 mus musculu	Q8utd7 human immun			O9hds4 aspergillus		_	Q37854 bacteriopha
SUMMARIES	ţ							Q726G0 052ZQ	ACRS					0		UMAN			2	ICJA													Q37854
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O6jc68 glycine max Q7xb03 zea may8 (m Q51594 plasmid col Q6uvk2 malus domes Q7x897 zea may8 (m Q712a6 sinorhizobi Q8giz6 lactobacill P13071 citrobacter P41495 sarcophaga P36641 mus mueculu Q49223 glycine max P83233 aloe vera (Q15894 homo sapien
Q6JC68 Q7XB03 Q51594 Q61594 Q6156 Q712A6 Q6GIZ6 B1OA CITFR TWOF_SARBU UF03 MOUSE O49223 VTIN ALOVR Q15894
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ALIGNMENTS

		Plasmodium.	ertea M., '', Koo H.L., dwell S.L., yum T.V., mmings L.M., Harris M.A., A.B., hite O.R.,	model rodent malaria rom an entry which is	gth 9; Indels 0; Gaps	nt).
9 AA.	ed) sequence update) annotation update)	yosiii yoelii. Alveolata; Apicomplexa; Haemosporida; Pla 13239; OM N.A.	STRAIN=17XNL; Pubmed=12546865; DOI=10.1038/nature01099; Carltcon J.M., Angiuoli S.V., Suh B.B., Kooji T.W., Pertea M., Silva J.C., Ermolaeva M.D., Allen J.E., Selengut J.D., Koo H.L., Stalva J.C., Ermolaeva M.D., Allen J.E., Shemay M.E., Bidwell S.L., Shallom S.J., van Aken S.B., Riedmuller S.B., Feldblyum T.V., Cho J.K., Quackenbush J., Sedegah M., Shoaibi A., Cummings L.M., Cuningham D.A., Preiser P.R. III, Raine J.D., Sinden R.B., Harris M.A., van Lin L.H., Janse C.J., Waters A.P., Smith H.O., White O.R., Salzberg S.L., Venter J.C., Fraser C.M., Hoffman S.L., Gardner M.	lysis of the is derived f hotgun (WGS)	DB 2; Len 1.6e+06; ches 0;	PRT; 9 AA. Created) Last sequence update) Last annotation update) regulatory chain (Fragment).
PRT;	26, Created) 26, Last sec 26, Last an	complexa;	038/natur V., Sub B V., Allen B., Alem Back D.S. B., Riedm Sedegah J, Raine P.R., Ber Waters A	i yoelii. shown hele genome 452.1;	Sco Pre 3;	PRT Creat Last Last regul
PRELIMINARY;	TrEMBLrel. TrEMBLrel. TrEMBLrel. protein.	Admuerty (1979); Plasmodium yoelii, Bukaryota; Alveolata; Api WCBI TaxID=73239; [1] SEQUENCE FROM N.A.	STRAIN=17XNL; Publed=12368865; DOI=10.1038/nature01099; Carlton J.M., Angiuoli S.V., Suh B.B., Ko Silva J.C., Ermolaeva M.D., Allen J.E., S Shuran J.D., Pop M., Kosack D. S., Shuran Shallom S.J., van Aken S.B., Riedmuller S Cho J.K., Quackenbush J., Sedegah M., Sho Florens L., Yates F.R. III, Raine J.D., S Cunningham D.A., Preiser P.R., Bergman L., van Lin L.H., Janse C.J., Waters A.P., Sm Salzberg S.L., Venter J.C., Fraser C.M., Carucci D.J.;	"Genome sequence and comparative ana parasite Plasmodium yoelii yoelii."; Nature 419:512-519(2002). Nature 419:512-519 (2002). EMBL/GenBank/DDBJ whole genome s preliminary data. EMBL, AABLO1005528; EAA19452.1; Hypothetical protein. SEQUENCE 9 AA; 1013 MW; 4684D447	Similarity 42.2%; Similarity 50.0%; 3; Conservative VLHDDL 6 : ::	PRELIMINARY; 6 (TrEMBLrel. 01, 6 (TrEMBLrel. 01, 8 (TrEMBLrel. 08, transcarbamoylase
RESULT 1 27RBX5 ID Q7RBX5	Q/KBX5; 01-MAR-2004 (01-MAR-2004 (01-MAR-2004 (Hypothetical	Name=F10/055; Plasmodium yoelii Eukaryota; Alveoli NCBI_TaxID=73239; [1] SEQUENCE_FROM N.A.	STRAIN=17XNL, Pubmed=123688 Carlton J.M., Silva J.C., E Peterson J.D., Cho J.K., Qua Florens L., Y Cunningham D., Van Lin L.H., Salzberg S.L., Salzberg S.L., Salzberg S.L., Salzberg S.L., Salzberg S.L., Salzberg S.L., Salzberg S.L.	Genome seque parasite Pla Nature 419:51 -!- CAUTION: EMBL/Gent prelimin EMBL; AABLOII Hypothetical SEQUENCE 9	Query Match Best Local Simi Matches 3; 1 VLH :	2 47556 1-NOV-199 1-NOV-199 1-NOV-199 spartate ame=pyrI;
RESULT Q7R8XS ID Q	A C C C C C C C C C C C C C C C C C C C	OC OC A A A A A A A A A A A A A A A A A	222222222222222	SOWER	o a a	RESULT Q47556 ID Q AC Q DT 0 DT 0 DT 0 DE A GN N

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01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
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                                                                                                                                        MEDLINE=83195078; PubMed=6302686;
Hoover T.A., Roof W.D., Foltermann K.F., O'Donovan G.A., Bencini D.A.
                                                                                                                                                                                                                                                                                   Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Gaps
                                                                                                                                                                        "Nucleotide sequence of the structural gene (pyrB) that encodes the catalytic polypeptide of aspartate transcarbamoylase of Escherichia
                                                                                                                                                                                                                                                                                                                                                                                                               01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-JUN-2003 (TrEMBLrel. 24, Last annotation update)
Alpha-myosin heavy chain (Fragment).
Oryctolagus cuniculus (Rabbit).
Eukaryota; Metazoa; Chordaea; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Lagomorpha; Leporidae; Oryctolagus.
                                                                         Pauza C.D., Karels M.J., Navre M., Schachman H.K.;
"Genes encoding Escherichia coli aspartate transcarbamoylase: the
           Bacteria; Proteobacteria; Gammaproteobacteria; Enterobacteriales;
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MEDLINE-84221901; PubMed-6328491;
Friedman D.J., Umeda P.K., Sinha A.M., Hsu H.-J., Jokovcic S.,
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                                                                                                                                                                                                                                      SEQUENCE 9 AA; 1085 MW; 99EFD723344AA1F1 CRC64;
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01-MAR-2004 (TrEMBLrel. 26, Last sequence update)
                                                                                               pyrB-pyrI operon.";
Proc. Natl. Acad. Sci. U.S.A. 79:4020-4024(1982)
                                                                                                                                                                                                     Proc. , Natl. Acad. Sci. U.S.A. 80:2462-2466 (1983)
EMBL; J01670; AAA24475.1; -.
NON TER
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      ventricular myosin heāvy chains.";
Proc. Natl. Acad. Sci. U.S.A. 81:3044-3048(1984)
EMBL; K01698; AAA31415.1; -.
PIR; 146868; 146868.
NON TER.
                                                                                                                                                                                                                                                           Score 19; DB 2; I
Pred. No. 1.6e+06;
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                                                               MEDLINE=82275057; PubMed=7051000;
              Enterobacteriaceae; Escherichia
NCBI_TaxID=83333;
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 Escherichia coli K12
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01-NOV-1996
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        Trypanosoma brucei.
Bukaryota; Buglenozoa; Kinetoplastida; Trypanosomatidae; Trypanosoma.
NCBI_TaxID=5691;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Gaps
                                                                                                                                                                      MEDLINE=95284106; PubMed=7766679; DOI=10.1016/0167-4838(95)00061-X;
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Acanthomorpha, Acanthopterygii, Percomorpha, Atherinomorpha,
Cyprinodontiformes, Aplocheilidae, Rivulinae, Gnatholebias.
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Homo sapiens (Human).
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
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                                                                                                                                                                                                    Hunt M., Koehler P.;
"Purification and characterization of phosphoenolpyruvate carboxykinase from Trypanosoma brucei.";
Biochim. Blophys. Acta 1249:15-22(1995).
PIR; S55696; S55696.
SEQUENCE 9 AA; 1063 MW; 35F2244331E05047 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                               1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                         Length 9;
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                                                                                                                                                                                                                                                                                                                                                                                                   7.8%; Score 17; DB 2; I
Local Similarity 33.3%; Pred. No. 1.6e+06;
Les 2; Conservative 3; Mismatches 1;
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Mol. Phylogenet. Evol. 13:289-301(1999)
EMBL; AF002591; AAD01074.1;
GO; GO:0005739; C:mitochondrion; IEA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            01-0CT-2003 (TrEMBLrel. 25, Created)
01-0CT-2003 (TrEMBLrel. 25, Last seq
01-0CT-2003 (TrEMBLrel. 25, Last ann
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Phosphoenolpyruvate carboxykinase.
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1 ILYQHLL 7
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Fri Jun

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

OM protein - protein search, using sw model

June Run on:

3, 2005, 07:10:14 ; Search time 85.5 Seconds (without alignments) 40.712 Million cell updates/sec

US-10-623-176-2 45 1 VLHDDLLEA 9 Title: Perfect score:

BLOSUM62 Scoring table:

Sequence:

2105692 seqs, 386760381 residues Gapop 10.0 , Gapext 0.5 Searched:

330156 Total number of hits satisfying chosen parameters:

seq length: 0 seq length: 9 Minimum DB Maximum DB Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

A_Geneseq_16Dec04:* 1: geneseqp1980s:* •• Database

geneseqp1980s:*

geneseqp1990s:* geneseqp2000s:* geneseqp2001s:*

geneseqp2002s:* geneseqp2003as:* geneseqp2003bs:*

geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

STIMMARTES

	Description	Aaw99196 Minor his	Aaw97375 HA-1 H-al	Adh40333 Human min	Aaw99197 Minor his	Aaw97374 HA-1 R-al	Adh40334 Human min	Aaw99195 Minor his	Aaw97572 T-cell ep	Aaw97373 Peptide e	Aay10122 T cell ep	MHC cla	Adk68732 Epitope 1		_	Adq10530 Cercopith	Abj20115 MHC bindi	Abu97233 Enzyme pe		Abu97234 Enzyme pe	Abu97231 Enzyme pe	2 Human s	Aae31275 Human mag	Adm96261 Human ser	Adm96258 Human ser	Abu97229 Enzyme pe
SUMMAKIES	ID	AAW99196	AAW97375	ADH40333	AAW99197	AAW97374	ADH40334	AAW99195	AAW97572	AAW97373	AAY10122	ABG79805	ADK68732	ADK05291	ADK05293	ADQ10530	ABJ20115	ABU97233	ABU97232	ABU97234	ABU97231	ADM96262	AAE31275	ADM96261	ADM96258	ABU97229
	DB	7	~	œ	~	~	œ	7	7	~	7	Ŋ	æ	8	æ	œ	9	9	9	9	9	œ	s	œ	œ	9
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æ	Query Match	100.0	100.0	100.0	82.2	82.2	82.2	80.0	80.0	66.7	62.2	62.2	62.2	62.2	62.2	62.2	60.09	57.8	57.8	57.8	57.8	57.8	57.8	57.8	57.8	55.6
	Score	45	45	45		37	37	36	36	30	28	28	28	28	28	28	27	56	56	26	26	26	56	26	26	25
	Result No.	п	7	e	4	S	9	7	80	σ	10	11	12	13		15	16	17	18	19	20	21	22	23	24	25

	Aaw39657 HPV16/18 Aaw4475 Peptide f Aay10561 HLA Class Abg80244 MHC class Adk69184 Epitope 1 Adk65290 Hepatitis
0440CUUU40UUU	9 2 AAN39657 9 2 AAN364759 . 9 2 AAN10561 9 5 ABGB0244 9 8 ADK69184
N N N N N 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	40 24 53.3 44 53.3 24

ALIGNMENTS

RESULT 1 AAW99196

AAW99196 standard; peptide; 9 AA

AAW99196;

(first entry) 20-MAY-1999 Minor histocompatibility antigen HA-1 T-cell epitope #2.

Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.

Homo sapiens

WO9905174-A1

04-FEB-1999.

98WO-NL000425. 23-JUL-1998;

97EP-00202303. 23-JUL-1997;

(UYLE-) RIJKSUNIV LEIDEN

Engelhard VH;

Hunt DF,

Goulmy EAJM,

WPI; 1999-153312/13.

A new minor histocompatibility antigen, HA-1 - useful to treat immune diseases and prevent rejection and host versus graft disease in bone marrow and organ transplantation.

Claim 3; Page 32; 47pp; English.

The present sequence represents a new peptide (Pl) constituting a T-cell epitope obtainable from the minor histocompatibility antigen HA-1. The peptide is immunogenic and can be used as part of a vaccine. Pl is used as a medicine, to induce tolerance for transplants, prevent rejection and/or graft versus host disease, or to treat (auto) immune diseases. In particular it can be used with bone marrow transplantation, in the treatment of severe aplastic anaemia, leukaemia, and immune deficiency

Sequence 9 AA;

Query Match

100.0%; Score 45; DB 2; Length 9;

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The invention relates to a novel method for providing epitopes of allelic variants of antigenic proteins from specific species based on single conclected polyworphism (SNP), by defining target protein/peptide or its subset, screening database of DNA encoding target protein, identifying, selecting allelic peptide/protein variants, expression product or its fragment encoded by DNA sequence having NP, creating variant epitopes, selecting epitopes binding to MHC protein. A protein of the invention has cytostatic activity, and may have a use in a vaccine. The method is cytostatic activity, and may have a use in a vaccine. The method is cytostatic selecting epitopes binding to MHC protein. A protein from the individuals, where the SNP profile of one or more individuals from a cancer. This is useful for diagnosing a disease in an individual by generating the SNP-related polymorphic profile. A method of the invention is useful for transplanting haematopoietic stem cells from a donor to a cepten progression, regression or onset of a treated disease. The present cells used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Providing allelic variant epitope of protein based on single nucleotide polymorphism by defining target protein, screening database of protein, identifying, selecting allelic variant protein, creating variant
                                                                           human, cytostatic, vaccine, SNP profile, cancer, leukaemia, minor histocomptibility antigen; mHAg; T cell epitope.
                                      Human minor histocomptibility antigen HA-1 T cell eptiope
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Disclosure; Page 82; 119pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          AAW99197 standard; peptide; 9 AA.
                                                                                                                                                                                                                                                                                                                13-JUN-2002; 2002EP-00013423.
                                                                                                                                                                                                                                                                        13-JUN-2003; 2003WO-EP006251.
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  11-MAR-2004 (first entry)
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                                                                                                                                               Homo sapiens.
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                                                                                                                                                                                                                                                                                                                                                                        Intron; minor histocompatibility antigen HA-1; typing allele; H allele; R allele; polymorphic nucleotide; HA-1 typing; bone marrow transplant; severe aplastic anaemia; leukaemia; immune deficiency disease; ss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Typing minor histocompatibility antigen HA-1 - by amplifying and identifying HA-1 R and HA-1 H alleles of HA-1, used for, e.g. detection
                       Gaps
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                       Indels
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  ; Pred. No. 1.8e+06; 0; Mismatches 0;
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                                                                                                                                                                                                           AAW97375 standard; protein; 9 AA.
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98EP-00870125.
100.08;
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                       9; Conservative
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                                                                                                                                                                                                                                                                                                                                    HA-1 H-allele sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            of genetic aberrances.
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es 9; Conserv
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    Best Local Similarity
Matches 9; Conser
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02-JUN-1998;
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Matches
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AAW97375
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                                                                               Gaps
                                                                            .;
0
                      Length 9;
                                                                            Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Minor histocompatibility antigen HA-1 T-cell epitope #3.
reacch 100.0%; Score 45; DB 8; Local Similarity 100.0%; Pred. No. 1.8e+06; Les 9; Conservative 0; Mismatches
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ADH40333 standard; peptide; 9 AA.

RESULT 3 ADH40333 ADH40333;

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CTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide;
FRATURE:
NAME/KEX: SITE
J. COGATION: (1)..(9)
US-10-623-176-2
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larity 100.0%; Pred. No. 1.3e+06;
Conservative 0; Mismatches 0;
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Best Local Similarity
Matches 9; Conserv
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                                                                                                  3, 2005, 07:48:11 ; Search time 67 Seconds (without alignments) 46.435 Million cell updates/sec
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Sequence 45,
Sequence 10,
Sequence 57,
Sequence 1, A
Sequence 1, A
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| cgn2 6/ptodata/1/pubpas/USO7 PUBCOMB.pep:*
| cgn2 6/ptodata/1/pubpas/USO6 NEW PUB.pep:*
| cgn2 6/ptodata/1/pubpas/USO8 PUBCOMB.pep:*
| cgn2 6/ptodata/1/pubpas/USO8 NEW PUB.pep:*
| cgn2 6/ptodata/1/pubpas/USO8 NEW PUB.pep:*
| cgn2 6/ptodata/1/pubpas/USO8 PUBCOMB.pep:*
              GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-10-791-217-2
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US-10-623-176-45
US-10-623-176-47
US-10-791-217-5
US-10-623-176-10
US-10-623-176-40
US-10-623-176-40
US-10-623-176-40
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                                                                          OM protein - protein search, using sw model
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Gapop 10.0 , Gapext 0.5
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Sequence 4, Appli
Sequence 15, Appl
Sequence 23, Appl
Sequence 28, Appl
Sequence 163, Appl
Sequence 163, Appl
Sequence 164, Appl
Sequence 166, Appl
Sequence 166, Appl
Sequence 167, Appl
Sequence 167, Appl
Sequence 3, Appli
Sequence 3, Appli
Sequence 3, Appli
Sequence 14, Appl
Sequence 16, Appl
Sequence 17, Appli
Sequence 18, Appli
Sequence 114, Appli
Sequence 112, Appli
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Sequence 2, Application US/10623176

Publication No. US20040092446A1

GENERAL INFORMATION:

APPLICANT: GOULMY, Els A.J.M.

APPLICANT: Engelhard, Victor H.

TITLE OF INVENTION: HA-1 epitopes and uses thereof

FILE REFERENCE: 2183-6040W

CURRENT APPLICATION HUMBER: US/10/623,176

FRIOR APPLICATION NUMBER: 09/489,760

PRIOR FILING DATE: 2000-01-21

PRIOR FILING DATE: 1997-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

NUMBER OF SEQ ID NOS: 77

SOSTWARE: PATENTIN VEV. 2.1
US-10-245-871-785
US-10-245-871-809
US-10-253-286-785
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US-09-834-765-132
US-09-834-765-349
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ORGANISM: Artificial Sequence
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Gaps

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Length 9; Indels

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OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
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OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
FEATURE:
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Publication No. US20040092446A1

GENERAL INFORMATION:
APPLICANT: Goulmy, Els A.J.M.
APPLICANT: Hunt, Donald F.
TILE REFERENCE: 2183-60470S

CURRENT APPLICATION NUMBER: US/10/623,176

CURRENT APPLICATION NUMBER: 09/489,760

PRIOR APPLICATION NUMBER: 09/489,760

PRIOR FILING DATE: 2003-07-18

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 2000-01-24

NUMBER OF SEQ ID NOS: 77
                                                                     GENERAL INFORMATION:
APPLICANT: Goulmy, Els A.J.M.
APPLICANT: Hunt, Donald F.
APPLICANT: Els Hart, Victor H.
TITLE OF INVENTION: HA-1 epitopes and uses thereof
FILE REPERENCE: 2183-60470S
CURRENT APPLICATION NUMBER: US/10/623,176
CURRENT PILING DATE: 2000-01-21
PRIOR APPLICATION NUMBER: 09/489,760
PRIOR APPLICATION NUMBER: EP 97202303.0
PRIOR PILING DATE: 1997-07-23
PRIOR PILING DATE: 1998-07-23
PRIOR FILING DATE: 1998-07-23
PRIOR FILING DATE: 1998-07-23
PRIOR FILING DATE: 1998-07-23
PRIOR FILING DATE: 2000-01-24
SROID RILING DATE: 2000-01-24
NUMBER OF SEQ ID NOS: 77
SOFTWARE: PARENTIN Ver. 2.1
Sequence 41, Application US/10623176
Publication No. US20040092446A1
GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                TYPE: PRT
ORGANISM: Artificial Sequence
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ORGANISM: Artificial Sequence
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Best Local Similarity
Matches 8; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           NAME/KEY: SITE
LOCATION: (1)..(9)
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US-10-623-176-45
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Publication No. US20050031612A1

GENERAL INFORMATION:

APPLICANT: Goulny, Elsa A.J.M.

TITLE OF INVENTION: Minor histocompatibility antigen HA-1: target antigen for immunot

TITLE OF INVENTION: Of tumors

FILE REFERENCE: 2183-6479US

CURRENT APPLICATION NUMBER: US/10/861,335

CURRENT APPLICATION NUMBER: PCT/NL02/00791

PRIOR APPLICATION NUMBER: EP 01204704.9

PRIOR APPLICATION NUMBER: EP 01204704.9

PRIOR FILING DATE: 2000-12-05

NUMBER OF SEQ ID NOS: 10

SEQ ID NO 1

LENGTH: 9
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Best Local Similarity 100.0%; Pred. No. 1.3e+06;
Matches 9; Conservative 0; Mismatches 0; Indels
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                                                                                                                                                                               APPLICANT: Goulmy, Elsa A.J.M
APPLICANT: Hunt, Donald F
APPLICANT: Hunt, Donald F
APPLICANT: Bogelhard, Victor H
TILE OF INVENTION: The HA-1 Antigen
FILE REPERENCE: 2183-4285US
CURRENT APPLICATION NUMBER: US/10/791,217
CURRENT APPLICATION NUMBER: US/09/489,760
PRIOR FILING DATE: 2000-01-21
PRIOR FILING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: Patentin Ver: 2.1
SENGTH. 9
LENGTH: 9
                                                                                                       Sequence 2, Application US/10791217 Publication No. US20040191268A1 GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ; OTHER INFORMATION: HA-1 peptide
US-10-861-335-1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   TYPE: PRT
ORGANISM: Artificial sequence
FEATURE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TYPE: PRT
ORGANISM: Artificial Sequence
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US-10-861-335-1
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US-09-269-250E-20
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Query Match
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29.211 Million cell updates/sec
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/cgn2_6/ptodata/1/iaa/PcTUS_COMB.pep:*
/cgn2_6/ptodata/1/iaa/PcTUS_COMB.pep:*
        GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-269-250E-18
US-09-269-250E-38
US-09-808-126-5
US-09-803-331A-3
US-08-803-331A-7
US-08-803-331A-7
US-08-803-331A-3
US-08-803-331A-3
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US-08-208-036-9
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Maximum Match 100%
Listing first 45 summaries
                                       OM protein - protein search, using sw model
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                                                                                       US-10-623-176-2
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Match Length
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Sequence 18, Application US/09269250E
Patent No. 6830883
GENERAL INFORMATION:
GENERAL INFORMATION:
APPLICANT: GOUINY, Elsa
TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REPRENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269, 250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: PatentIn version 3.1
SEQ ID NO 18
LENGTH: 9
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TITLE OF INVENTION: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REFERENCE: 56994
CURRENT APPLICATION NUMBER: US/09/269,250E
CURRENT FILING DATE: 1999-05-21
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
SEQ ID NO 20
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100.0%; Pred. No. 4.1e+05;
US-09-586-472-20
US-09-528-706-20
US-08-399-411-75
US-08-999-411-75
US-09-419-826-1
US-09-419-826-1
US-09-586-472-75
US-09-586-472-75
US-09-586-706-75
US-08-343-602-2
US-08-459-568-72
US-08-459-568-72
US-08-459-568-72
US-08-459-568-72
US-08-399-411-72
US-08-399-411-72
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US-08-399-411-72
US-08-556-76
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ORGANISM: Artificial Sequence
FEATURE:
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ORGANISM: Artificial Sequence
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Best Local Similarity 100.
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Length

DB 4;

Score 37;

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APPLICANT: KOYama, Tanetoshi
APPLICANT: Sato, Yoshihiro
TITLE OF INVENTION: DECAPRENYL DIPHOSPHATE SYNTHETASE GENE
NUMBER OF SEQUENCES: 31
CORRESPONDENCE ADDRESS:
ADDRESSEE: KENYON & KENYON
STREET: 1500 K Street. N W
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Nishino, Tokuzo
Koyama, Tanetoshi
Sato, Yoshihiro
TITLE OF INVENTION: DECAPRENYL DIPHOSPHATE SYNTHETASE GENE
NUMBER OF SEQUENCES: 31
CORRESPONDENCE ADDRESS:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               MEDIUM TYPE: Floppy disk
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
                                                                                                                                                                                                                                                                                                              COMPUTER: IBM PC compatible
COMPUTER: IBM PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS
SOFTWARE: Patentin Release #1.0, Version #1.30
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/025,819
FILING DATE: 19-FEB-1998
CLASSIFICATION: 435
PRIOR APPLICATION DATA:
APPLICATION NUMBER: JP 251675
FILING DATE: 17-SEP-1997
ATTORNEY/AGENT INFORMATION:
NAME: KRAIJIIAA, HOURT:
NAME: KRAIJIIAA, HOURT:
NAME: CRAIJIIAA, HOURT:
NAME: CRAILILIAN NUMBER: 39,546
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              APPLICATION NUMBER: US/09/808,126
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   STREET: 1500 K Street, N.W CITY: Washington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  ADDRESSEE: KENYON & KENYON
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Sequence 5, Application US/09808126
Patent No. 6410280
GENERAL INFORMATION:
APPLICANT: Obata, Shusei
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-220-4200
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CURRENT APPLICATION DATA:
                                                                                                                                                                                                                                                                             COMPUTER READABLE FORM:
MEDIUM TYPE: Floppy disk
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COMPUTER READABLE FORM:
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Best Local Similarity 80.0
Matches 4; Conservative
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US-09-808-126-5
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                                                                                                                                                                                                                                           Squence 29, Application US/09269250E

Facent No. 6830883
GENERAL INFORMATION:
APPLICANT GOULDY, Elsa
TITLE OF INVENTON: METHOD FOR TYPING OF MINOR HISTOCOMPATIBILITY ANTIGEN HA-1
FILE REFERENCE: 58994
CURRENT APPLICATION NUMBER: US/09/269,250E
NUMBER OF SEQ ID NOS: 38
SOFTWARE: Patentin version 3.1
LENGTH: 9
LENGTH: 9
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Sequence 38, Application US/09269250E

Sequence 38, Application US/09269250E

Sequence 38, Application US/09269250E

GENERAL INFORMATION:

APPLICANT: Goulmy, Elsa

TITLE OF INVENTION:

FILE REPERENCE: 58994

CURRENT APPLICATION NUMBER: US/09/269,250E

CURRENT FILING DATE: 1999-05-21

NUMBER OF SEQ ID NOS: 38

SOFTWARE: Patentin version 3.1

SEQ ID NO 38

LENGTH: 8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    ; LOCATION: (3)...(3)
; OTHER INFORMATION: Xaa represents a histidine (H) or an arginine (R) residue
US-09-269-250E-29
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                                                   Pred. No. 4.1e+05;
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88.9%; Pred. No. 4.1e+05;
tive 0; Mismatches 1;
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APPLICANT: Obata, Shuse; APPLICANT: Nishino, Tokuzo
                                                                      88.9%;
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Best Local Similarity 86.>
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1 VLRDDLLEA 9
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                                                                      Best Local Similarity
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Matches 8; Conser
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ORGANISM: HUMAN
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US-09-025-819-5
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C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Date: 15-Feb-1997 #sequence_revision 13-Mar-1997 #text_change 17-Mar-1999
C;Accession: $68004
R;Edgar, P.F.
R;Edgar, P.F.
PEBS Lett. 375, 159-161, 1995
A;Title: Hucolin, a new corticosteroid-binding protein from human plasma with structural A;Reference number: $68004; MUID:96087107; PMID:7498469
ferredoxin a2 - Ja
conopressin S - Co
chlorophyll a/b-bi
dextraneric protein
glycine cleavage s
transaldolase (EC
neuropeptide Grb-A
neuropeptide Grb-A
calliPMRPamide 10
gamma subunit of P
protein-glutamine
neuropeptide Antho
hypothetical protein
glycoprotein compo
ribosomal protein
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   S69165
BB28495
BB398002
BB39801
S66419
S66419
B57253
B57444
C574444
C574444
C574444
A44787
A4478
A4788
A
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      RESULT 1
      868004
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3, 2005, 07:30:19 ; Search time 16.25 Seconds (without alignments) 53.289 Million cell updates/sec OM protein - protein search, using sw model US-10-623-176-10 1 VLRDDLLEA 9 June Title: Perfect score:

Run on:

GenCore version 5.1.6 Copyright (c) 1993 - 2005 Compugen Ltd.

283416 seqs, 96216763 residues BLOSUM62 Gapop 10.0 , Gapext 0.5 Searched:

Scoring table:

Sequence:

Total number of hits satisfying chosen parameters:

791

Minimum DB seq length: 0 Maximum DB seq length: 9

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

pir2:* pir3:* pir4:* PIR_79:* |: pir1:*

Database

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

A;Accession: S68004 A;Status: preliminary A;Molecule type: protein A;Residues: 1-7 <EDG>

		Description	hucolin, 75K chain	hypothetical prote	protein kinase C i	elastase - Pseudom	macrophage cytotox	T-cell receptor be	biotin A - Citroba	T-cell receptor be	T-cell receptor be	-	T-cell receptor be	acetylcholinestera	calsequestrin, fas	T-cell receptor be	T-cell receptor be	T-cell receptor be	. T-cell receptor be		late G1-69 protein	rece	T-cell receptor be	sucrose 3-glucosyl		major protein anti	phosphoglycerate t	pilE protein - Esc	T-cell receptor be	chitin-binding pro	158K exoantigen -
	£	1	9			S20446	A60427	PH0942	140697	PT0679	PT0601	B35640	PT0533	A34026	B39040	PT0628	PT0722	PT0576	PT0557	: A61328	PH0108	PT0562	B30572	A39841	A37027	D60274	: T11779	S25266	PT0602	242	B33099
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dip	Query		œ	38.1	35.7	33.3	33.3	31.0	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	28.6	26.2	26.2	26.2	26.2	26.2	26.2
	2	SCOLE	16	16	15	14	14	13	12	12	12	12	12	12	12	12	12	12	12	12	12	12	12	12	12	11	11	11	11	11	11
,	Result		7	7	æ	4	S.	9	7	80	σ	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	56	27	28	29

Query Match 38.1%; Score 16; DB 2; Length 7; Best Local Similarity 100.0%; Pred. No. 2.8e+05; Matches 3; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
6y 4 ppr 6
Db 4 DDL 6
RESULT 2
PC4131
hypotherical protein 8 [imported] - Pseudomonas aeruginosa (fragment)
C.Decise: Febucomorias actuginosa C.Date: 20-Abr-2000 #sequence revision 20-Abr-2000 #text change 09-Jul-2004
C; Accession: PC4131
R;Kawasaki, S.; Arai, H.; Igarashi, Y.; Kodama, T.
Gene 167, 87-91, 1995
A, Title: Sequencing and characterization of the downstream region of the genes encoding r
y for biosynthesis of heme d1.
A; Reference number: JC4552; MUID: 96144254; PMID: 8566817
A;Accession: PC4131
A;Status: preliminary
A; Molecule type: DNA
A;Residues: 1-8 <kaw></kaw>
A; Cross-references: UNIFROT: P95412; DDBJ: D50473; NID: g1217594
A;Note: this ORF is not annotated in GenBank entry PSENIRC, release 113.0

ö Similarity 100.0%; Pred. No. 2.8e+05; 3; Conservative 0; Mismatches 0; Indels 9 DDF Query Match Best Local S: Matches 3 4 8

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Gaps

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RESULT

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Gaps

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C;Species: Rattus norvegicus (Norway rat)
C;Daces: 09-0ct-1992 #sequence_revision 09-0ct-1992 #text_change 30-May-1997
C;Daces: 09-0ct-1992 #sequence_revision 09-0ct-1992 #text_change 30-May-1997
C;Accession: PH0942
R;Gold, D.P.; Offiner, H.; Sun, D.; Wiley, S.; Vandenbark, A.A.; Wilson, D.B.
J. Exp. Med. 174, 1467-1475, 1991
A;Fitle: Analysis of T cell receptor beta chains in Lewis rats with experimental allergic A;Reference number: PH0891; MUID:92078857; PMID:1836012
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Diotin A - Citrobacter freundii (fragment)
C;Species: Citrobacter freundii
C;Accession: I40697
R;Shiuan, D.; Campbell, A.
Gene 67, 203-211, 1988
A;Title: Transcriptional regulation and gene arrangement of Escherichia coli, Citrobacter A;Reference number: I40697; MUID:89006280; PMID:2971595
A;Reference number: I40697
A;Reference number: I40697
A;Reference problemary; translated from GB/EMBL/DDBJ
A;Molecule type: DNA
A;Residues: 1-4 cRES>
A;Residues: 1-4 cRES>
A;Residues: 1-4 cRES>
A;Cross-references: UNIPROT:P13071; GB:M21922; NID:g144434
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C;Species: Mus musculus (house mouse)
C;Species: IJ-ul-1992 #sequence_revision 17-Jul-1992 #text_change 30-May-1997
C;Accession: PT0679; PT0708
B;Feeney, A.J.
J. Exp. Med. 174, 115-124, 1991
A;Title: Junctional sequences of fetal T cell receptor beta chains have few N regions. A;Reference number: PT0509; MUID:91277601; PMID:1711558
                                                                                                                                                                                                                                                                                                                                    A;Accession: PH0942
A;Molecule type: mRNA
A;Residues: 1-9 <GOL>
A;Residues: 19 <GOL>
A;Experimental source: complete Freund's adjuvant-immunized lymph node
A;Note: the authors translated the codon TGC for residue 2 as Ala
C;Keywords: T-cell receptor
                                                                                         T-cell receptor beta chain V-D-J region (clone 13) - rat (fragment)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    A;Experimental source: day 18 fetal thymus, strain BALB/c, 154-2J
A;Accession: PT0708
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31.0%; Score 13; DB 2; Length 9;
Best Local Similarity 100.0%; Pred. No. 2.8e+05;
Matches 3; Conservative 0; Mismatches 0; Indels
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A;Experimental source: newborn thymus, strain BALB/c, 161-2B
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     28.6%; Score 12; DB 2; Le
ilarity 100.0%; Pred. No. 2.8e+05;
Conservative 0; Mismatches 0;
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R;Jones, C.M.; Prince, C.A.; Williams, J.S.
R;Jones, C.M.; Prince, C.A.; Williams, J.S.
Exp. Hematol. 19, 704-709, 1991
A;Title: Purification and amino acid analysis of a human macrophage cytotoxicity-inducin A;Reference number: A60427; WUID:91372335; PMID:1909970
A;Accession: A60427
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    R,Kessler, E.; Safrin, M.; Peretz, M.; Burstein, Y.
FBBS Lett. 299, 291-293, 1992
A;Title: Identification of cleavage sites involved in proteolytic processing of Pseudomc
A;Reference number: S20446; MUID:92183956; PMID:1544509
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               A;Cross-references: UNIPROT:Q7M4R5
A;Note: the sequence from the text on page 706 is inconsistent with that from page 708
C;Keywords: cytokine
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ö
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                      C;Species: Rattus norvegicus
C;Date: 25-Aug-2003 #sequence_revision 25-Aug-2003 #text_change 25-Aug-2003
C;Date: 25-Aug-2003 #sequence_revision 25-Aug-2003 #text_change 25-Aug-2003
C;Accession: A59489
A;Reference number: A59489
A;Reference number: A59489
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  elastase - Pseudomonas aeruginosa
C;Species: Pseudomonas aeruginosa
C;Date: 19-Mar_1997 #sequence_revision 24-Jul-1997 #text_change 24-Jul-1997
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C;Species: Homo sapiens (man)
C;Date: 12-Feb-1993 #sequence_revision 12-Feb-1993 #text_change 09-Jul-2004
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Pred. No. 2.8e+05;
2; Mismatches 0; Indels
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Pred. No. 2.8e+05;
0; Mismatches 1; Indels
                                                                                                                                                                                                                                                                    A;Status: preliminary
A;Molecule type: protein
A;Residues: 1-7 <NEG>
A;Experimental Source: strain Wistar, liver
A;Note: p-Hydroxyacetophenone-Sepharose binding protein
- rat (fragment)
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Best Local Similarity 50.0%;
Matches 2; Conservative ;
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ilarity 75.0%;
Conservative
rotein kinase C inhibitor
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A;Molecule type: protein
A;Residues: 1-7 <KES>
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Gaps

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GenCore version 5.1.6
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June Run on:

3, 2005, 07:11:54; Search time 77.5 Seconds (without alignments) 59.467 Million cell updates/sec

US-10-623-176-10 42 1 VLRDDLLEA 9

Title: Perfect score: Sequence:

Scoring table:

BLOSUM62 Gapop 10.0 , Gapext 0.5

1612378 seqs, 512079187 residues Searched:

1455 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 9

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database

UniProt_03:*
1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

	1	Description	Q7xb03 zea mays (m	zea n	homo	homo	Q8lpt5 zea mays (m	P82568 streptococc	P72279 rhodococcus	Q80h91 newcastle d	Q9ue26 homo sapien				Q51594 plasmid col				_	_	Q8utd7 human immun	Q85710 rous sarcom			Q6jc68 glycine max	Q93sr0 staphylococ	Q9qvj8 mus sp. mep		simian	simian	simian	simian	Q6qsx9 simian viru
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SUMMARIES			03	P7	42	P6	TS TS	89	79	91	92	¥0	39	74	94	69	35	20	22		70	01	97	54		80	18	55	25	70	60	E1	63
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	9	a :	~	~	~	7	~	7	~	7	N	~	7	~	~	~	7	~	~	7	~	~	~	~	7	~	7	7	~	~	7	~	0
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	9	score	18	18	16	16	16	16	15	15	15	15	15	14	14	14	14	14	14	14	14	14	13	13	13	13	13	13	13	13	13	13	13
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Q64ix4 fundulus he P82992 rhodopseudo	_	Q61cv2 homo sapien	Q6uvk2 malus domes	_	Q9pykl simian viru	Q64ix3 fundulus he	P13071 citrobacter	099182 gnatholebia	049223 glycine max	P83233 aloe vera (Q6u7r2 cryptococcu	Q7z4n9 homo sapien
264IX4 IPYR RHOVI	27RY89	26LCV2	26UVK2	Q91BM8	29PYK1	2641X3	BIOA CITFR	0991 <u>8</u> 2	049223	JIIN ALOVR	26U7 <u>R</u> 2	2724N9
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ALIGNMENTS

PRELIMINARY; PRT; 8 AA. ; 2003 (TrEMBLrel. 25, Created) -2003 (TrEMBLrel. 25, Last sequence update)	(TrEMBLrel. 25, Last nthase 2 (Fragment). aize). Viridiplantae; Streptc ta; Magnoliophyta; Lil e; Panicoideae; Androg	[1] - Seguence FROM N.A. STRAIN-Y-3; MEDLINE=22779048; PubMed=12897253; MEDLINE=22779048; PubMed=12897253; MEDLINE=22779048; PubMed=12897253; "Contrasting effects of selection on sequence diversity and linkage disequilibrium at two phytoene synthase loci."; Plant Cell 15:1795-1806(2003). Plant Cell 15:1795-1806(2003). Plant Cell 15:1795-1806(2003). Plant Cell 15:1795-1806(2003). NON TER 8 8 SEQÜENCE 8 AA; 915 MW; 5D1862CAB4072054 CRC64;	ch 42.9%; Score 18; DB 2; Length 8; 1 Similarity 75.0%; Pred. No. 1.6e+06; 3; Conservative 1; Mismatches 0; Indels 0; Gaps 0; 1 VLRD 4 2 ILRD 5	7 2 QYX8P7 PRELIMINARY; PRT; 9 AA. QYX8P7; 9 AA. QYX8P7; 0. Created) 10CCT-2003 (TrEMBLrel. 25, Last sequence update) 05JUL-2004 (TrEMBLrel. 27, Last annotation update) Name=psy2; 05. Cas mays (Maize). Sea mays (Maize). Each mays (Maize). Sebraryota; Waridiplantae; Streptophyta; Embryophyta; Tracheophyta; Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae; PACCAD clade; Panicoideae; Andropogoneae; Zea. [1] SEQUENCE FROM N.A. SEQUENCE FROM N.A. STRAIN=Y-13, and W-16;
RESULT 1 Q7XB03 ID Q7XB03 AC Q7XB03, DT 01-OCT-2003 DT 01-OCT-2003		RN [1] RP SEQUENCE FROM RC STRAIN=Y-3; RX MEDLINE=22775 RA Palaisa K.A.; RT "Contrasting RT "contrasting RT disequilibriu RL Plant Cell 15 DR EMBL; AY30061 PT NON TER FT NON TER SQ SEQUENCE 8	Query Match Best Local Si Matches 3 Qy 1 1 Db 2 2	RESULT 2 07X8P7 D 07X8P7 AC 07X8P7 DT 01-0CT-2003 (TD 01-0CT-2003 (TD 01-0CT-2003 (TD 01-0CT-2003 (TD 01-0CT-2004 (TD 01-0CT-2

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Query Match
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DEVVDA 9
  [2]
SEQUENCE FROM N.A.
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                                                                                                 SEQUENCE FROM N.A.
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                                                                   expression
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SEQUENCE
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P82568;
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Pinte S., Guerardel C., Deltour S., Godwin A.K., Leprince D.;
Pinte S., Guerardel C., Deltour S., Godwin A.K., Leprince D.;
Identification of a second G-C-rich promoter conserved in the human,
murine and rat tumor suppressor genes HICL.";
Oncogene 23:4023-4031(2004).
EMBL; AJ550616; CAD79467.1; -.
                                                                                                                                  Gaps
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Palaisa K.A., Morgante M., Williams M., Rafalski A.;
"Contrasting effects of selection on sequence diversity and linkage disequilibrium at two phytoene synthase loci.";
Plant Cell 15:1795-1806(2003).
EMBL, AX300641; AAP55379.1; -.
                                                                                                                                                                                                                                                                                                               Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
NCBI_TaxID=9606;
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Mammalia, Butheria, Primates, Catarrhini, Hominidae, Homo.
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MEDLINE=90005483; PubMed=2792104;
Wignal A., Rahuel C., El-Maliki B., London J., Le Vankim C.,
Blanchard C.D.;
"Molecular Analysis of glycophorin A and B gene structure and
expression ...; ";
Eur. J. Blochem. 184:337-344(1989).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1; Indels
                                                                                                                                0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    38.1%; Score 16; DB 2; Length 9; 60.0%; Pred. No. 1.6e+06; tive 1; Mismatches 1; Indels
                                                                                                            Length 9;
                                                                                      9 AA; 1016 MW; 1D1C62CAB4072054 CRC64;
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05-JUL-2004 (TrEMBLrel. 27, Last sequence update)
05-JUL-2004 (TrEMBLrel. 27, Last annotation update)
M1.V glycoprotein (exon P3B) (Fragment).
Homo sapiens (Human).
                                                                                                                                                                                                                                                              Last sequence update)
Last annotation update)
                                                                                                                                                                                                                                                                                                                                                                                                                                                SEQUENCE 9 AA; 964 MW; 5B5E6DDB1681AAA7 CRC64;
                                                                                                         Score 18; DB 2; I
Pred. No. 1.6e+06;
1; Mismatches 0;
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75.0%;
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Best Local Similarity
3, Conserve
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DTMEA 7
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NON TER
SEQUENCE
                                                                                                                                                                                                                                                                                             Name=HIC1
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Matches
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Q7KYP6
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Ching A.S., Caldwell K.S., Jung M., Dolan M., Smith O.S., Tingey S., Morgante M., Rafalski J.A.;
Morgante M., Rafalski J.A.;
Submitted (APR-2002) to the EMBL/GenBank/DDBJ databases.
EMBL; AY094310; AAM21836.1; --
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Spermatophyta, Magnoliophyta, Liliopsida, Poales, Poaceae,
PACCAD clade, Panicoideae, Andropogoneae, Zea.
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                                                                                       "Molecular Analysis of glycophorin A and B gene structure and
TISSUE=Miltenberger class V;
Vignal A., Rahuel C., El-Maliki B., London J., Le Vankim
Blanchard C.D.;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    0; Indels
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                                                                                                                                                                                                                                                    Cartron J.D.; Submitted (JUL-1989) to the EMBL/GenBank/DDBJ databases. EMBL; X15824; CAA33822.1; -. NON TER 9
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Bacteria; Firmicutes; Lactobacillales; Streptococcaceae;
                                                                                                                                                                                                                                                                                                                                           NON TER 9 9 SEQÜENCE 9 AA; 1073 MW; 50D145B7244AB403 CRC64;
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01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-OCT-2003 (TrEMBLrel. 25, Last annotation update)
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Last annotation update)
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75.0%; Pred. No. 1.6e+06;
:ive 1; Mismatches 0;
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01-0CT-2002 (TrEMBLrel. 22, Last seque
01-0CT-2002 (TrEMBLrel. 22, Last anno
Beta-expansin-like protein (Fragment)
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                                                                                                                          expression . . . . ";
Eur. J. Biochem. 184:0-0(1989)
                                                                                                                                                                                                                                TISSUE=Miltenberger class V;
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Best Local Similarity 75.v.
3; Conservative
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GenCore version 5.1.6 -
Copyright (c) 1993 - 2005 Compugen Ltd.
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- protein search, using sw model OM protein June Run on:

3, 2005, 07:10:14 ; Search time 85.5 Seconds (without alignments) 40.712 Million cell updates/sec

US-10-623-176-10 Title:

1 VLRDDLLEA 9 Perfect score: Sequence:

Gapop 10.0 , Gapext 0.5 BLOSUM62 Scoring table:

2105692 seqs, 386760381 residues Searched:

330156 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 9

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

A_Geneseq_16Dec04: .. Database

geneseqp1980s:* geneseqp1990s:*

geneseqp2003as:* geneseqp2000s:* geneseqp2001s:* geneseqp2002s:*

geneseqp20048:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

uo	Minor his	HA-1 R-al	Human min	Minor his	HA-1 H-al	Human min	Minor his	T-cell ep	Prenyl di	Human MHC	Zif268 zi	C trachom	Chlamydia	Human MHC	Human can	Peptide e	Rabbit pe	Peptide #	Rat Rb-in	Mammalia					
Description		Aaw97374	Adh40334	Aaw99196	Aaw97375	Adh40333	Aaw99195	Aaw97572	Aaw47427	Aau71428	Aab81550	Adm18256	Adm18218	Aau71211	Abr19981	Abr19426	Abr18988	Abr19578	Abr19174	Abr19780	Aaw97373	Adf72091	Aab12035	Aau80810	Ade84629
ID	AAW99197	AAW97374	ADH40334	AAW99196	AAW97375	ADH40333	AAW99195	AAW97572	AAW47427	AAU71428	AAB81550	ADM18256	ADM18218	AAU71211	ABR19981	ABR19426	ABR18988	ABR19578	ABR19174	ABR19780	AAW97373	ADF72091	AAB12035	AAU80810	ADE84629
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Score	42	42	42	37	37	37	36	36	27	25	24	24	24	23	23	23	23	23	23	23	22	22	21	21	21
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Abp48411 Zinc fing Abp48419 Zinc fing Abp48419 Zinc fing Abg79371 CXCR zinc Abp96179 Zinc fing Ada62245 Zinc fing Ada62245 Zinc fing Adm20444 Synthetic Adm20441 Synthetic Adm20431 Synthetic Adm20431 Synthetic Adm30431 Synt	Aay80162 HLA-A2 re Aab12092 Ad40 cel
7. S ABP48411 7 5 ABP48414 7 5 ABP48419 7 5 ABP79371 7 6 ABP96179 7 7 ADA62240 7 7 ADA62245 7 7 ADM20444 7 7 ADM20441 7 7 ADM20441 7 7 ADM2048 8 2 AAW26557 8 2 AAW26557 8 4 AAB46568 8 4 AAB46535 8 6 ABP99038	9 3 AAY80162 9 3 AAB12092
221 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	21 50.0 21 50.0
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ALIGNMENTS

Minor histocompatibility antigen HA-1 T-cell epitope #3 AAW99197 standard; peptide; 9 AA. (first entry) 20-MAY-1999 AAW99197; RESULT 1 AAW99197

Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.

Homo sapiens

WO9905174-A1

04-FEB-1999.

98WO-NL000425. 23-JUL-1998; 97EP-00202303. 23-JUL-1997;

(UYLE-) RIJKSUNIV LEIDEN

Engelhard VH; Hunt DF, Goulmy EAJM,

WPI; 1999-153312/13.

A new minor histocompatibility antigen, HA-1 - useful to treat immune diseases and prevent rejection and host versus graft disease in bone marrow and organ transplantation.

Disclosure; Page 15; 47pp; English.

The present sequence represents a new peptide (PI) constituting a T-cell peptiope obtainable from the minor histocompatibility antigen HA-1. The peptide is immunogenic and can be used as part of a vaccine. Pl is used as a medicine, to induce tolerance for transplants, prevent rejection and/or graft versus host disease, or to treat (auto) immune diseases. In particular it can be used with bone marrow transplantation, in the treatment of severe aplastic anaemia, leukaemia, and immune deficiency

Sequence 9 AA;

Query Match

Length 9; DB 2; 100.0%; Score 42;

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The invention relates to a novel method for providing epitopes of allelic variants of antigenic proteins from specific species based on single nucleotide polyworphism (SNP), by defining target protein/peptide or its subset, screening database of DNA encoding target protein, identifying, selecting allelic peptide/protein variants, expression product or its fragment encoded by DNA sequence having SNP, creating variant epitopes, selecting epitopes binding to MMC protein. A protein of the invention has useful for generating as SNP profile of one or more individuals from a given species by applying the method for several protein from the individuals, where the SNP profile was related to disease, preferably cancer. This is useful for dispnosing a disease in an individual by generating the SNP-related polymorphic profile. A method of the invention is useful for transplanting haematopoietic stem cells from a donor to a copient and treating encer, preferably leuksemia, and for determining the progression, regression or onset of a treated disease. The present sequence is used in the exemplification of the invention.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Providing allelic variant epitope of protein based on single nucleotide polymorphism by defining target protein, screening database of protein, identifying, selecting allelic variant protein, creating variant
                                                                                   human, cytostatic; vaccine; SNP profile; cancer; leukaemia; minor histocomptibility antigen; mHAg; T cell epitope.
                                            Human minor histocomptibility antigen HA-1 T cell eptiope.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Disclosure; Page 82; 119pp; English
                                                                                                                                                                                                                                                                                                                                       13-JUN-2002; 2002EP-00013423.
                                                                                                                                                                                                                                                                                             L3-JUN-2003; 2003WO-EP006251
11-MAR-2004 (first entry)
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nes 9; Conser
                                                                                                                                                                                                     WO2003106692-A2.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     The present sequence represents part of the minor histocompatibility antigen HA-1 R-allele. The specification describes methods for typing alleles (preferably the H and R alleles) of the minor histocompatibility antigen HA-1 in a sample, which comprise detecting polymorphic nucleotides in the cDNA or genomic nucleic acids of the alleles. The methods can be used for HA-1 typing for bone marrow transplants, severe aplastic anaemia, leukaemia and immune deficiency diseases, as well as detection of genetic aberrances. The probes and primers of the invention can be used to screen for the HA-1 alleles. The HA-1 peptides can be used anti-idiotypic B cells and/or T cells and antibodies
                                                                                                                                                                                                                                                                                                                                                                                                     Intron, minor histocompatibility antigen HA-1; typing allele; H allele; R allele; polymorphic nucleotide; HA-1 typing; bone marrow transplant; severe aplastic anaemia; leukaemia; immune deficiency disease; ss.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Typing minor histocompatibility antigen HA-1 - by amplifying and identifying HA-1 R and HA-1 H alleles of HA-1, used for, e.g. detection
                         Gaps
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  Pred. No. 1.8e+06;
                       0; Mismatches
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98EP-00870125.
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  100.08;
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  Best Local Similarity
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Les 9, Conserv
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Matches
                       Matches
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                                                                                                                                                                                                                                                                                Minor histocompatibility antigen; HA-1; T-cell epitope; immunological; graft versus host disease; bone marrow transplant; leukaemia; vaccine; diagnosis; aplastic anaemia; immune deficiency disease.
                               Gaps
                             ö
Length 9;
                            Indels
                                                                                                                                                                                                                                                         Minor histocompatibility antigen HA-1 T-cell epitope #2
100.0%; Score 42; DB 8; I
100.0%; Pred. No. 1.8e+06;
                               0; Mismatches
                                                                                                                                                                    AAW99196 standard; peptide; 9 AA.
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ADH40334 standard; peptide; 9 AA.

RESULT 3 ADH40334

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ADH40334;

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Fri Jun
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3, 2005, 07:48:11; Search time 67 Seconds (without alignments) 46.435 Million cell updates/sec
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1: /cgn2_6/ptodata/1/pubpaa/USO7_PUBCOMB.pep:*
2: /cgn2_6/ptodata/1/pubpaa/PCT_MRP_PUB.pep:*
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11: /cgn2_6/ptodata/1/pubpaa/USO9_PUBCOMB.pep:*
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13: /cgn2_6/ptodata/1/pubpaa/USO9_NEW_PUB.pep:*
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19: /cgn2_6/ptodata/1/pubpaa/USO0_PUBCOMB.pep:*
GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    1465611 segs, 345679903 residues
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                                                                                                                                            OM protein - protein search, using sw model
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Gapop 10.0 , Gapext 0.5
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42
1 VLRDDLLEA 9
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                                                                                                                                                                                                                      June
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Perfect score:
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

		Appl	Appli	Appl	Appl	Appli	Appli	Appli	Appli	Appli	Appl	Appl	Appl	Appl
	c	10,	'n		46,		7	ì	ť	ì	48	23,	41,	45
	Description	Seguence 10,	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence	Sequence 41,	Segmence
	QI	US-10-623-176-10	US-10-791-217-5	US-10-623-176-42	US-10-623-176-46	US-10-623-176-2	US-10-791-217-2	US-10-861-335-1	US-10-623-176-1	US-10-791-217-1	US-10-623-176-48	US-10-623-176-23	US-10-623-176-41	US-10-623-176-45
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0 0 0 72 4 4	349 339	Sequence 4, Appli Sequence 15, Appl Sequence 132, App Sequence 3, Appli	44.00		216, 6, 7, 219, 565,	Sequence 54, Appl Sequence 109, App Sequence 219, App Sequence 565, App	1001104
5 US-10-623-17 5 US-10-623-17 5 US-10-623-17	US-09-834-765- US-10-623-176 US-10-271-708	15 15 15 15 15 15	0-623-1 0-791-2 9-990-1	US-09-990-186-2 US-09-990-186-2 US-09-989-994-2 US-09-989-994-2	US-09-989-994 US-10-122-675 US-09-935-430 US-09-935-430	US-09- US-10- US-10- US-10-	10-280-340-56 -10-057-475B-1 -10-057-475B-1 -10-154-884B-1 -10-154-884B-1
29 69. 29 69. 28 66.	25 25 25 25 25 25 25 25 25 25 25 25 25 2	20 24 57.1 21 23 54.8 22 23 54.8 23 23 54.8	22 2 2 2 2 2 3 3 3 3 3 3 3 3 3 3 3 3 3	2222	221212	22222	40 21 50.0 41 21 50.0 43 21 50.0 44 21 50.0 45 21 50.0

ALIGNMENTS

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FEATURE:
OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
FEATURE:
NAME/KEX: SITE
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US-10-623-176-10

Sequence 10, Application US/10623176

Publication No US20040092446A1

GENERAL INPORMATION:

APPLICANT: Goulmy, Bls A.J.M.

APPLICANT: Engelhard, Victor H.

TITLE OF INVENTION: HA-1 epitopes and uses thereof

FILE REFERENCE: 2183-6047US

CURRENT APPLICATION NUMBER: US/10/623,176

CURRENT FILING DATE: 2003-07-18

PRIOR APPLICATION NUMBER: 09/489,760

PRIOR FILING DATE: 2000-01-21

PRIOR FILING DATE: 1997-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 2000-01-21

SEQIEND NOS: 77

SEQIEND NO 10
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             TYPE: PRT
ORGANISM: Artificial Sequence
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US-10-623-176-10
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Gaps

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Query Match
Best Local Similarity 100.
Matches 8; Conservative
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nes 8; Conservative
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2 VLRDDLLE
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; LOCATION: (1)..(9)
US-10-623-176-46
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            OTHER INFORMATION: Description of Artificial Sequence: HA-1 peptide
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100.0%; Score 42; DB 16; Length 9;
Best Local Similarity 100.0%; Pred. No. 1.3e+06;
Matches 9; Conservative 0; Mismatches 0; Indels
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Sequence 42, Application US/10623176

Publication No. US20040092446A1

GENERAL INFORMATION:

APPLICANT: Goulmy, Els A.J.M.

APPLICANT: Engelhard, Victor H.

TITLE OF INVENTION: HA-1 epitopes and uses thereof FILE REFERENCE: 2183-60470S

CURRENT APPLICATION HWHER: US/10/623,176

CURRENT FILING DATE: 2003-07-18

PRIOR FILING DATE: 2000-01-21

PRIOR FILING DATE: 1997-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-23

PRIOR FILING DATE: 1998-07-24

PRIOR FILING DATE: 1998-07-24

NUMBER OF SEQ ID NOS: 77

SEQ ID NO 42

LENGTH: 9

LENGTH: 9

LENGTH: 9
                                                                                                                                                           GENERAL INFORMATION:
FUDILCATION NO. US20040191268A1
GENERAL INFORMATION:
APPLICANT: Goulmy, Elsa A.J.M
APPLICANT: Hunt, Donald F
APPLICANT: Engelhard, Victor H
ITILE OF INVENTION: The HA-1 Antigen
FILE REFERENCE: 2183-4285US
CURRENT APPLICATION NUMBER: US/10/791,217
CURRENT APPLICATION NUMBER: US/09/489,760
PRIOR PTLING DATE: 2000-01-21
PRIOR APPLICATION NUMBER: PCT/NL98/00424
PRIOR PTLING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: PATENTING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: PATENTING DATE: 1998-07-23
NUMBER OF SEQ ID NOS: 17
SOFTWARE: PATENTING DATE: 1998-07-23
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ORGANISM: Artificial Sequence
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ORGANISM: Artificial sequence
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LOCATION: (1)..(9)

US-10-623-176-42
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29.211 Million cell updates/sec
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6: /cgn2 6/ptodata/1/iaa/PCTUS_COMB.pep:*
GenCore version 5.1.6
Copyright (c) 1993 - 2005 Compugen Ltd.
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US-09-269-250E-29

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US-09-269-250E-29

US-09-269-250E-29

US-08-819-14

US-08-856-411-20

US-08-856-411-20

US-08-816-829-20

US-08-816-829-20

US-08-816-829-20

US-08-816-829-20

US-08-816-829-75

US-08-819-811-75

US-08-819-811-75

US-08-819-811-75

US-08-819-811-72

US-08-816-819-72

US-08-816-819-73
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                                                                                                                                                                                                                                                                                                                                                              513545 segs, 74649064 residues
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Maximum Match 100%
Listing first 45 summaries
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Gapop 10.0 , Gapext 0.5
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42
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Match Length DB
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No.
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Sequence 72, Appl Sequence 18, Appl Sequence 11, Appl Sequence 11, Appl Sequence 11, Appl Sequence 11, Appl Sequence 50, Appl Sequence 51, Appl Sequence 51, Appl Sequence 1186, Ap Sequence 1186, Ap Sequence 1186, Ap Sequence 1189, Ap Sequence 1189, Ap Sequence 20, Appl Sequence 20, Appl		MINOR HISTOCOMPATIBILITY ANTIGEN HA-1 DE .cial Sequence: Exon Fragments	ngth 9; Indels 0; Gaps 0;	OMPATIBILITY ANTIGEN HA-1
9 4 US-09-528-706-72 9 4 US-09-528-706-78 6 3 US-09-025-819-11 6 4 US-09-808-126-11 6 4 US-09-803-951-11 8 3 US-08-747-599A-51 8 3 US-08-747-599A-51 8 3 US-08-747-599A-51 9 4 US-09-839-54186 7 3 US-09-187-859-1186 7 4 US-09-839-5428-1186 8 3 US-09-187-859-1189 8 3 US-09-187-859-1189 8 4 US-09-839-5428-189 9 1 US-08-441-4178-20 9 2 US-08-444-4178-20	ALIGNMENTS	/09269250E POR TYPING OF: US/09/269,25G -05-21 3.1	100.0%; Score 42; DB 4; Length 100.0%; Pred. No. 4.1e+05; vative 0; Mismatches 0; Inde 9	09-269-250B-20 equence 20, Application US/09269250E equence 20, Application US/09269250E EXERTAL INFORMATION: APPLICANT: GOULMY, E1sa APPLICANT: GOULMY, E1sa APPLICANT: GOULMY, E1s9 FILE REFERENCE: 58994 CURRENT APPLICATION NUMBER: US/09/269,250E CURRENT FILING DATE: 1999-05-21 NUMBER OF SEQ ID NOS: 38 SOFTWARR: Patentin version 3.1 LENGTH: 9 TYPE: RRT FEATURE: FRT FEATURE: CONTRANTION: Description of Artificial Sequence: Exon Fraggr
211 221 201 200 200 200 200 200 200 200		250E- 18, 0.068 10NONT: C NT: C F INN FEREN APPI APPI APPI APPI C 18 C 18 C 18 C 18 C 18 C 18 C 18 C 18	ttch 9, Conser 1 VLRDDLLEA	250E-20 20, Applicat o. 683083; INTFORMATION; F INVENTION; FFIRENCE; 589 FFILING DATE; OF SEQ ID NOS E: Patentin v O 20 E: Patentin v O 20 E: Patentin v E:
2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		RESULT 1 Sequence 18 ; Sequence 18 ; Patent No.; Patent P.;	Query Match Best Local Batches Matches Dy 1	RESULT 2 US-09-269-256 Sequence 20 Sequenc

Length 9;

88.1%; Score 37; DB 4;

Query Match

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39,048
                                                                                                                                                                             ; TOPOLOGY: linear
; MOLECULE TYPE: peptide
US-09-217-609A-11
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MOLECULE TYPE: peptide
    REGISTRATION NUMBER:
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STATE:
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                                                                                                                                                                                      Sequence 29, Application US/09269250E

| Sequence 29, Application US/09269250E
| Patent No. 6830683
| GENERAL INFORMATION:
| APPLICANT: GOULD, BESALD, BESALD, BETLICANT: GOULD, BETLICANT: GOULD, BETLICANT: SAPPLICANT: 58994
| CURRENT APPLICANTION: UNMBER: US/09/269,250E
| CURRENT FILING DATE: 1999-05-21
| NUMBER OF SEQ ID NOS: 38
| SEQ ID NO 29
| LENGTH: 9
| LENGTH: 9
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1. LOCATION: (3) ... (3)

1. OCHER INORWATION: Xaa represents a histidine (H) or an arginine (R) residue

US-09-2508-259-2508-29
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APPLICANT: KOIKE, Ayumi
APPLICANT: KOIKE, Ayumi
APPLICANT: KOIKE, Ayumi
APPLICANT: KOYAMA, Kyozo
APPLICANT: SHINIY Tanetoshi
APPLICANT: SHINIY Wasto
APPLICANT: SHINIY Wasto
APPLICANT: CHO, Yenwin
TITLE OF INVENITON: Prenyl Diphosphate Synthetase Genes
NUMBER OF SEQUENCES: 29
CORRESPONDENCE ADDRESS:
ADDRESSEE: Kenyon & Kenyon
STREET: 1025 Connecticut Avenue, NW - Suite 600
CITY: Washington
STREE: DC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Query Match 85.7%; Score 36; DB 4; Length 9; Best Local Similarity 88.9%; Pred. No. 4.1e+05; Matches 8; Conservative 0; Mismatches 1; Indels
                       1; Indels
    Pred. No. 4.1e+05;
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MEDIUM TYPE: 3.5 inch Floppy disk
COMPUTER: 18 PC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS 6.2
SOFTWARE: WordPerfect 6.1 for Windows
CURRENT APPLICATION DATA:
APPLICATION NUMBER: US/09/217,609A
FILING DATE:
                       0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          US-09-217-609A-11
; Sequence 11, Application US/09217609A
Patent No. 6071733
; GENERAL INFORMATION:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            CLASSIFICATION:
PRIOR APPLICATION DATA:
APPLICATION NUMBER: 08/873,235
FILING DATE: 11-Jun-1997
ATTORNEY/AGENT INFORMATION:
NAME: TOPFENETTI, Judith L.
88.98;
  Best Local Similarity 88.9
Matches 8; Conservative
                                                           1 VLRDDLLEA 9
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1 VLHDDLLEA 9
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                                                                                                                                                                                                                                                                                                                                                                                                                                                    TYPE: PRT
ORGANISM: Human
FEATURE:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  APPLICANT: MURAMATSU, Masayoshi
APPLICANT: KOIKE, Ayumi
APPLICANT: CGURA, Kyozo
APPLICANT: KOYAMA, Tanetcshi
APPLICANT: SHIMIZU, Naoto
APPLICANT: SHIMIZU, Naoto
APPLICANT: PHIMIZU, Premyl Diphosphate Synthetase Genes
NUMBER OF SEQUENCES: 29
CORRESPONDENCE ADDRESS:
ADDRESSEE: Kenvon F V-
                                                                                                                                                                                                                                                                          Score 27; DB 3; Length 9;
Pred. No. 4.1e+05;
3; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Query Match

64.3%; Score 27; DB 3; Length 9;
Best Local Similarity 57.1%; Pred. No. 4.1e+05;
Matches 4; Conservative 3; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SSEE: Kenyon & Kenyon
: 1025 Connecticut Avenue, NW - Suite 600
Washington
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           COMPUTER READBLE FORM:
MEDIUM TYPE: 3.5 inch Floppy disk
COMPUTER: TBM FC compatible
OPERATING SYSTEM: PC-DOS/MS-DOS 6.2
SOFTWARE: WordPerfect 6.1 for Windows
CURENT APPLICATION DATA:
APPLICATION NUMBER: US/08/873,235B
FILING DATE: 11-Jun-1997
CLASSIFICATION: 435
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PRIOR APPLICATION DATA:

APPLICATION NUMBER: JP 15441/1996

FILING DATE: 14-Jun-1996

ATTORNEY/AGENT INFORMATION:

NAME: TOFFENETT, Judith L.

REGISTRATION NUMBER: 39,048

REFERENCE/DOCKET NUMBER: 10235/1

TELECOMMUNICATION INFORMATION:

TELEPHONE: 202-429-1776
    10235/1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Sequence 11, Application US/08873235B Patent No. 6174715
REFERENCE/DOCKET NUMBER: 102
TELECOMMUNICATION INFORMATION:
TELEPHONE: 202-429-1776
TELEFAX: 202-429-0796
INFORMATION FOR SEQ ID NO: 11:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                                 64.3%;
57.1%;
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INFORMATION FOR SEQ ID NO:
SEQUENCE CHARACTERISTICS:
                                                                                                                                                                                                                                                                               Query Match 64.3
Best Local Similarity 57.1
Matches 4; Conservative
                                                                                                                                            9 amino acids
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